I U C L I D

Data s e t

Existing Chemical

CAS No.

CAS Name

ID: 13752-51-7 13752-51-7

 ${\tt N-oxydiethylenethiocarbamyl-N-oxydiethylenesulfenamide}$

237-335-9 EINECS No.

Producer Related Part

Company:

Creation date:

09-NOV-2000

Substance Related Part

Company:

Creation date:

09-NOV-2000

Rubber and Plastic Additives (RAPA) HPV Panel Memo:

Printing date:

Revision date:

12-OCT-2001

12-OCT-2001 Date of last Update:

28 Number of Pages:

Chapter: 1, 2, 3, 4, 5, 7 Chapter (profile):

Reliability (profile): Reliability: without reliability, 1, 2, 3, 4

Flags: without flag, confidential, non confidential, WGK Flags (profile): (DE), TA-Luft (DE), Material Safety Dataset, Risk

Assessment, Directive 67/548/EEC, SIDS

Date: 12-OCT-2001 1. General Information ID: 13752-51-7

1.0.1 OECD and Company Information

Type: lead organisation

American Chemistry Council (formerly Chemical Manufacturers Name:

Association) Rubber and Plastic Additives (RAPA) HPV Panel

Street: 1300 Wilson Boulevard Town: 22209 Arlington, VA

United States Country: Phone: 703-741-5600 Telefax: 703-741-6091

11-OCT-2001

Type: cooperating company Name: Bayer Corporation United States Country:

11-OCT-2001

Type: cooperating company

Ciba Specialty Chemicals Corporation Name:

United States Country:

11-OCT-2001

Type: cooperating company Name: Crompton Corporation

United States Country:

11-OCT-2001

cooperating company Type: Name: Flexsys America L.P.

United States Country:

11-OCT-2001

Type: cooperating company

Noveon, Inc (formerly BF Goodrich) Name:

United States Country:

11-OCT-2001

cooperating company Type:

R.T. Vanderbilt Company, Inc. Name:

Country: United States

11-OCT-2001

Type: cooperating company

The Goodyear Tire & Rubber Company Name:

United States Country:

11-OCT-2001

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Date: 12-OCT-2001

1. General Information ID: 13752-51-7

Type: cooperating company
Name: The Lubrizol Corporation

Country: United States

11-OCT-2001

Type: cooperating company

Name: UOP, LLC. Country: United States

11-OCT-2001

1.0.2 Location of Production Site

-

1.0.3 Identity of Recipients

_

1.1 General Substance Information

Substance type: organic Physical status: solid

Purity: 95 - 99 % w/w

25-APR-2001

1.1.0 Details on Template

_

1.1.1 Spectra

-

1.2 Synonyms

Morpholine, 4-[(morpholinothiocarbonyl)thio]-25-APR-2001

Cure-Rite® 18 25-APR-2001

Good-Rite® 3030x18

25-APR-2001

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Date: 12-OCT-2001

1. General Information ID: 13752-51-7

1.3 Impurities

CAS-No: 729-46-4

EINECS-No:

EINECS-Name: Dimorpholine Thiuram Disulfide

Contents: < 5 % w/w

25-APR-2001

CAS-No: 34986-62-4

EINECS-No:

EINECS-Name: [4-(4'-morpholinodithion) thioxomethyl-morpholine]

Contents: < .5 % w/w

25-APR-2001

CAS-No: 110-91-8
EINECS-No: 203-815-1
EINECS-Name: morpholine
Contents: < .02 % w/w

25-APR-2001

CAS-No: 59-89-2

EINECS-No:

EINECS-Name: N-nitrosomorpholine

Contents: < .005 % w/w

25-APR-2001

1.4 Additives

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1.5 Quantity

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1.6.1 Labelling

_

1.6.2 Classification

_

1.7 Use Pattern

_

1.7.1 Technology Production/Use

_

1.8 Occupational Exposure Limit Values

-

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Date: 12-OCT-2001 1. General Information ID: 13752-51-7

1.9 Source of Exposure

1.10.1 Recommendations/Precautionary Measures

1.10.2 Emergency Measures

1.11 Packaging

1.12 Possib. of Rendering Subst. Harmless

1.13 Statements Concerning Waste

1.14.1 Water Pollution

1.14.2 Major Accident Hazards

1.14.3 Air Pollution

1.15 Additional Remarks

1.16 Last Literature Search

1.17 Reviews

1.18 Listings e.g. Chemical Inventories

- 4/28 -

2. Physico-chemical Data

2.1 Melting Point

Value: 124.3 degree C

Method: other: (calculated) MPBPWIN (v1.31)

Year: 1999 GLP: no

Testsubstance: other TS: molecular structure

Comparable to BFG MSDS data of >/= 132 °C Remark:

Melting Point: 251.84 deg C (Adapted Joback Method) Result: Melting Point: 92.44 deg C (Gold and Ogle Method) Mean Melt Pt: 172.14 deg C (Joback; Gold,Ogle Methods) Selected MP: 124.32 deg C (Weighted Value)

(2) valid with restrictions Reliability:

Accepted calculation method

Flaq: Critical study for SIDS endpoint

11-OCT-2001 (1)

2.2 Boiling Point

Value: 353 degree C

Method: other: (calculated) MPBPWIN (v1.31) - Adapted Stein and Brown

Method

1999 Year: GLP:

Testsubstance: other TS: molecular structure

Remark: BFGoodrich MSDS indicates Not Applicable

Reliability: (2) valid with restrictions

Accepted calculation method

Flaq: Critical study for SIDS endpoint

11-OCT-2001 (1)

2.3 Density

Type: density Value: .6 g/cm3

Method: other: historical data Testsubstance: as prescribed by 1.1 - 1.4

25-APR-2001 (2)

2.3.1 Granulometry

- 5/28 -

2.4 Vapour Pressure

Value: .0000153 hPa at 25 degree C

Method: other (calculated): MPBPWIN (v1.31)

Year: 1999 GLP: no

Testsubstance: other TS: molecular structure

Result: Vapor Pressure Estimations (25 deg C):

(Using BP: 352.97 deg C (estimated))

(Using MP: 124.32 deg C (estimated))

Using MP: 124.32 deg C (estimated))

VP: 5.17E-006 mm Hg (Antoine Method)

VP: 1.15E-005 mm Hg (Modified Grain Method)

VP: 2.32E-005 mm Hg (Mackay Method)

Selected VP: 1.15E-005 mm Hg (Modified Grain Method)

Reliability: (2) valid with restrictions
Accepted calculation method

Flag: Critical study for SIDS endpoint

11-OCT-2001 (1)

2.5 Partition Coefficient

log Pow: -.84

Method: other (calculated): KOWWIN Program (v1.65)

Year: 1999 GLP: no

Testsubstance: other TS: molecular structure
Reliability: (2) valid with restrictions
Accepted calculation method

Flag: Critical study for SIDS endpoint

11-OCT-2001 (1)

2.6.1 Water Solubility

Value: 62.85 g/l at 25 degree C

Method: other: (calculated) WSKOW (v1.36)

Year: 1999 GLP: no

Testsubstance: other TS: molecular structure
Result: Log Kow (estimated) : -0.84

Log Kow (experimental): not available from database Log Kow used by Water solubility estimates: -0.84

Equation Used to Make Water Sol estimate:

Log S (mol/L) = 0.796 - 0.854 log Kow - 0.00728 MW + Correction (used when Melting Point NOT available)

Correction(s): Value
----Amine, aliphatic 1.008
Multi-Nitrogen Type -1.310

Log Water Solubility (in moles/L): -0.597 Water Solubility at 25 deg C (mg/L): 6.285e+004

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Date: 12-OCT-2001 ID: 13752-51-7

2. Physico-chemical Data

Reliability: (2) valid with restrictions Accepted calculation method

Critical study for SIDS endpoint Flag:

11-OCT-2001 (1)

2.6.2 Surface Tension

2.7 Flash Point

2.8 Auto Flammability

Value: 275 degree C
Remark: Self-Ignition Temperature
25-ADR-2001

25-APR-2001 (2)

2.9 Flammability

2.10 Explosive Properties

2.11 Oxidizing Properties

2.12 Additional Remarks

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3.1.1 Photodegradation

air Type: INDIRECT PHOTOLYSIS Sensitizer: OH

Conc. of sens.: 1560000 molecule/cm3

Rate constant: .000000000002156 cm3/(molecule * sec)

Degradation: 50 % after .6 hour(s)

other (calculated): AOP (v1.89): Method: 1999 Year:

Test substance: other TS: molecular structure Reliability: (2) valid with restrictions Accepted calculation method

Flaq: Critical study for SIDS endpoint

11-OCT-2001 (1)

3.1.2 Stability in Water

See IUCLID data sets on CAS# 95-31-8; 102-77-2; 95-33-0; 4979-32-2

3.1.3 Stability in Soil

3.2 Monitoring Data (Environment)

3.3.1 Transport between Environmental Compartments

fugacity model level III Type:

Media: other: air, water, soil, sediment

Air (Level I): Water (Level I): Soil (Level I): Biota (L.II/III): Soil (L.II/III):

other: EPIWIN Level III Fugacity Model Method:

Year: 1999

Result: Media Concentration Half-Life Emissions Fugacity (percent) (hr) (kg/hr) (atm) 0.00657 1.19 1000 1.47e-013 Air 900 50.2 1000 9.26e-015 Water 1000 Soil 49.7 900 3.38e-013 Sediment 0.0927 8.53e-015 3.6e + 003

Persistence Time: 763 hr

Reliability:

Reaction Time: 1.24e+003 hr Advection Time: 1.99e+003 hr

Percent Advected: 38.4 (2) valid with restrictions

Percent Reacted: 61.6

Accepted calculation method Flag: Critical study for SIDS endpoint

- 8/28 -

3. Environmental Fate and Pathways

11-OCT-2001 (1)

3.3.2 Distribution

3.4 Mode of Degradation in Actual Use

3.5 Biodegradation

See IUCLID data sets on CAS# 95-31-8; 102-77-2; 95-33-0; 4979-32-2

3.6 BOD5, COD or BOD5/COD Ratio

3.7 Bioaccumulation

Species: other: calculation

Exposure period: Concentration:

3.16 BCF:

Elimination:

Method: other: BCF Program (v2.13)

Year: 1999 GLP: no

Test substance: other TS: molecular structure Result: Log Kow (estimated) : -0.84

Log Kow (experimental): not available from database

Log Kow used by BCF estimates: -0.84

Equation Used to Make BCF estimate:

Log BCF = 0.50

Correction(s): Correction Factors Not Used for Log Kow < 1</pre>

Estimated Log BCF = 0.500 (BCF = 3.162)

Accepted calculation method

11-OCT-2001 (1)

3.8 Additional Remarks

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AQUATIC ORGANISMS

4.1 Acute/Prolonged Toxicity to Fish

See also IUCLID data sets on CAS# 95-31-8; 102-77-2; 95-33-0; 4979-32-2

Type: other: calculation

Species: other: Fish
Exposure period: 96 hour(s)

Unit: g/l Analytical monitoring: no

LC50: 86.036

Method: other: ECOSAR v0.99e

Year: 1999 GLP: no

Test substance: other TS: molecular structure

Remark: Chemical may not be soluble enough to measure this predicted

effect.

Reliability: (2) valid with restrictions

Accepted calculation method

Flag: Critical study for SIDS endpoint

11-OCT-2001 (1)

Type: other: calculation Species: other: Saltwater Fish

Exposure period: 96 hour(s)

Unit: g/l Analytical monitoring: no

LC50: 4.992

Method: other: ECOSAR v0.99e

Year: 1999 GLP: no

Test substance: other TS: molecular structure

Remark: Chemical may not be soluble enough to measure this predicted

effect.

Reliability: (2) valid with restrictions

Accepted calculation method

Flag: Critical study for SIDS endpoint

11-OCT-2001 (1)

Type: other: calculation

Species: other: Fish

Exposure period: 14 day

Unit: g/l Analytical monitoring: no

LC50: 99.248

Method: other: ECOSAR v0.99e

Year: 1999 GLP: no

Test substance: other TS: molecular structure

Remark: Chemical may not be soluble enough to measure this predicted

(1)

effect.

Reliability: (2) valid with restrictions
Accepted calculation method

11-OCT-2001

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4.2 Acute Toxicity to Aquatic Invertebrates

See also IUCLID data sets on CAS# 95-31-8; 102-77-2; 95-33-0; 4979-32-2

Type: other: calculation
Species: Daphnia sp. (Crustacea)

Exposure period: 48 hour(s)

Unit: g/l Analytical monitoring: no

LC50: 75.767

Method: other: ECOSAR v0.99e

Year: 1999 GLP: no

Test substance: other TS: molecular structure

Remark: Chemical may not be soluble enough to measure this predicted

effect.

Reliability: (2) valid with restrictions
Flag: Critical study for SIDS endpoint

11-OCT-2001 (1)

Type: other: calculation

Species: Mysidopsis bahia (Crustacea)

Exposure period: 96 hour(s)

Unit: q/l Analytical monitoring: no

LC50: 188

Method: other: ECOSAR v0.99e

Year: 1999 GLP: no

Test substance: other TS: molecular structure

Remark: Chemical may not be soluble enough to measure this predicted

effect.

Reliability: (2) valid with restrictions

Accepted calculation method

Flag: Critical study for SIDS endpoint

11-OCT-2001 (1)

Type: other: calculation
Species: Daphnia sp. (Crustacea)

Exposure period: 16 day

Unit: mg/l Analytical monitoring: no

EC50: 1121

Method: other: ECOSAR v0.99e

Year: 1999 GLP: no

Test substance: other TS: molecular structure

Remark: Chemical may not be soluble enough to measure this predicted

effect.

Reliability: (2) valid with restrictions

Accepted calculation method

11-OCT-2001 (1)

- 11/28 -

4.3 Toxicity to Aquatic Plants e.g. Algae

See also IUCLID data sets on CAS# 95-31-8; 102-77-2; 95-33-0; 4979-32-2

Species: other algae: green algae

Endpoint: growth rate
Exposure period: 96 hour(s)

Unit: g/l Analytical monitoring: no

EC50: 40.223 ChV: 0.779

Method: other: ECOSAR v0.99e

Year: GLP: no

Test substance: other TS: molecular structure

Remark: Chemical may not be soluble enough to measure this predicted

effect.

Reliability: (2) valid with restrictions
Accepted calculation method

Flag: Critical study for SIDS endpoint

11-OCT-2001 (1)

4.4 Toxicity to Microorganisms e.g. Bacteria

_

4.5 Chronic Toxicity to Aquatic Organisms

4.5.1 Chronic Toxicity to Fish

Species: other: fish

Endpoint: other Exposure period: 30 day

Unit: mg/l Analytical monitoring: no

ChV: 7012

Method: other: ECOSAR v0.99e

Year: 1999 GLP: no

Test substance: other TS: molecular structure

Remark: Chemical may not be soluble enough to measure this predicted

effect.

Reliability: (2) valid with restrictions

Accepted calculation method

11-OCT-2001 (1)

4.5.2 Chronic Toxicity to Aquatic Invertebrates

-

- 12/28 -

Date: 12-OCT-2001 ID: 13752-51-7 4. Ecotoxicity

TERRESTRIAL ORGANISMS

4.6.1 Toxicity to Soil Dwelling Organisms

other: calculation

Species: Eisenia fetida (Worm (Annelida), soil dwelling)
Endpoint: mortality

Exposure period: 14 day other: ppm Unit: LC50: 11449

other: ECOSAR v0.99e Method:

Year: 1999 GLP: no

Test substance: other TS: molecular structure

Remark: Chemical may not be soluble enough to measure this predicted

effect.

Reliability: (2) valid with restrictions Accepted calculation method

11-OCT-2001 (1)

4.6.2 Toxicity to Terrestrial Plants

4.6.3 Toxicity to other Non-Mamm. Terrestrial Species

4.7 Biological Effects Monitoring

4.8 Biotransformation and Kinetics

4.9 Additional Remarks

- 13/28 -

5.1 Acute Toxicity

5.1.1 Acute Oral Toxicity

Type: LD50 Species: rat

Strain: Sprague-Dawley

Sex:
Number of
 Animals:
Vehicle:

Value: 5200 mg/kg bw

Method: other: 40CFR Part 163.81-1

Year: GLP: yes

Test substance: other TS: Cure-Rite® 18, purity: not noted Remark: Rat/CD® Sprague-Dawley (Charles River)
Discriminating dose: LD0 = 2,700 mg/kg

Reliability: (1) valid without restriction

GLP study, Meets National standards method

Flag: Critical study for SIDS endpoint

11-OCT-2001 (3)

Type: LD50
Species: mouse
Strain: CD-1

Sex: male/female

Number of Animals: Vehicle:

Value: 9000 mg/kg bw

Method: other: 40CFR Part 163.81-1

Year: GLP: yes
Test substance: other TS: Cure-Rite® 18; purity: not noted

Remark: Diagram noting dogs: LDO = 4.050 mg/kg

Remark: Discriminating dose: LD0 = 4,050 mg/kg
LD50 (95% conf. Limits) = 11,000 mg/kg (5,100-16,900 mg/kg)
for males and 7,000 mg/kg (4,900-9,100 mg/kg) for females

Reliability: (1) valid without restriction

GLP study, Meets National standards method

Flag: Critical study for SIDS endpoint

11-OCT-2001 (4)

Type: LD50 Species: rat

Strain:
Sex:
Number of
Animals:
Vehicle:

Value: 5110 mg/kg bw

Method: other: Federal Hazardous Substances Act (Revised, Fed. Reg.,

September 17, 1964

Year: 1964 GLP: no

Test substance: other TS: Good-Rite® 3030x18 (Cure-Rite® 18); purity: not

 ${\tt noted}$

Remark: Discriminating dose: LD0 = 1,000 mg/kg

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Reliability: (2) valid with restrictions

25-APR-2001 (5)

Type: LD50 Species: rat

Strain: Sprague-Dawley

Sex:
Number of
 Animals:
Vehicle:

Value: 5000 mg/kg bw

Method: other: 40CFR Part 163.81-1

Year: GLP: yes

Test substance: other TS: Cure-Rite® 18; purity: not noted Remark: Rat/CD® Sprague-Dawley (Taconic Farms)

Discriminating dose: LD0 = 2,700 mg/kg

Reliability: (1) valid without restriction

GLP study, Meets National standards method

11-OCT-2001 (3)

5.1.2 Acute Inhalation Toxicity

Type: LC0 Species: rat

Strain: Sex: Number of Animals: Vehicle:

Exposure time: 1 hour(s)
Value: 164.4 mg/l

Method: other: Federal Hazardous Substances Act (Revised, Fed. Req.,

September 17, 1964)

Year: 1964 GLP: no

Test substance: other TS: Good-Rite® 3030x18 (Cure-Rite® 18); purity: not

noted

Remark: Test substances measured not analysed.

Reliability: (1) valid without restriction

Meets National standards method

Flag: Critical study for SIDS endpoint

11-OCT-2001 (5)

- 15/28 -

5.1.3 Acute Dermal Toxicity

Type: LD50 Species: rabbit

Strain:
Sex:
Number of
 Animals:
Vehicle:

Value: > 10000 mg/kg bw

Method: other:) Federal Hazardous Substances Act (Revised, Fed. Reg.,

September 17, 1964)

Year: 1964 GLP: no

Test substance: other TS: Good-Rite® 3030x18 (Cure-Rite® 18); purity: not

noted

Reliability: (1) valid without restriction Meets National standards method

Flag: Critical study for SIDS endpoint

11-OCT-2001 (5)

5.1.4 Acute Toxicity, other Routes

-

5.2 Corrosiveness and Irritation

5.2.1 Skin Irritation

-

5.2.2 Eye Irritation

Species: rabbit

Concentration:

Dose:

Exposure Time:
Comment:
Number of
Animals:

Result: not irritating

EC classificat.:

Method: other: Section 1500.42, Federal Hazardous Substances Act

Regulations, CFR 16, p. 125

Year: GLP: yes

Test substance: other TS: Cure-Rite® 18; purity: not noted

Result: No positive scores; one animal had scores of "1" for redness

and chemosis at day one.

Reliability: (1) valid without restriction

GLP study, Meets National standards method

11-OCT-2001 (6)

- 16/28 -

Species: rabbit

Concentration:

Dose:

Exposure Time: Comment: Number of Animals:

Result: irritating

EC classificat.:

Method: other: Federal Hazardous Substances Act (Revised, Fed. Req.,

September 17, 1964)

Year: 1964 GLP: no

Test substance: other TS: Good-Rite® 3030x18 (Cure-Rite® 18); purity: not

noted

Reliability: (1) valid without restriction
Meets National standards method

11-OCT-2001 (5)

5.3 Sensitization

-

5.4 Repeated Dose Toxicity

Species: rat Sex: male/female

Strain: Sprague-Dawley Route of admin.: oral feed Exposure period: 2 years

Frequency of

treatment: continuous daily

Post. obs.

period: none

Doses: 0, 20, 60, 200, or 600 ppm Control Group: yes, concurrent no treatment

NOAEL: 200 ppm LOAEL: 600 ppm Method: EPA OPP 82-5

Year: GLP: yes

Test substance: other TS: Commercial Cure-Rite® 18; purity: 96.8%

Result: A compound related increase in urothelial tumors, kidney

weights, non-neoplastic urinary tract abnormalities, and rales was observed in the high dose (600 ppm) males and females. Body weights also were significantly lower in the high dose in the high dose males and females. No compound-related effects on hematology, clinical chemistry, or urinalysis were noted.

Reliability: (1) valid without restriction

GLP guideline study

Flag: Critical study for SIDS endpoint

11-OCT-2001 (7)

- 17/28 -

5.5 Genetic Toxicity 'in Vitro'

Type: Bacterial reverse mutation assay

System of

testing: Salmonella typhimurium strains TA-1535, TA-1537, TA-1538,

TA-98, TA-100

Concentration: 0.5 to 1,000 ug/plate

Cytotoxic Conc.: With metabolic activation: 0.5 to 100 ug/plate (little to no

toxicity)

Without metabolic activation: 0.5 to 100 ug/plate (little to

no toxicity)

Metabolic

activation: with and without

Result: negative

Method: other: according to other: according to Ames et al (1975)

Mutation Res. 31:347-364; McCann et al. (1975) Proc. Nat.

Acad. Sci. 72:5135-5139

Year: 1975 GLP: no data

Test substance: other TS: Commercial Cure-Rite® 18 (Purified); purity: 97.5%

Remark: Signed QA assurance statement provided

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Flag: Critical study for SIDS endpoint

12-OCT-2001 (8) (9)

Type: Bacterial reverse mutation assay

System of

testing: Escherichia coli strain WP2urvA-

Concentration: 0.5 to 1,000 ug/plate

Cytotoxic Conc.: With metabolic activation: 0.5 to 100 ug/plate (little to no

toxicity)

Without metabolic activation: 0.5 to 100 ug/plate (little to

no toxicity)

Metabolic

activation: with and without

Result: negative

Method: other: according to Hinderer, R.K., B. Myhr, D.R. Jagannath,

S.M. Galloway, S.W. Mann, J.C.Riddle, and D.J. Brusick (1983)

Environ. Mutagen. 5:193-215

Year: 1983 GLP: no data

Test substance: other TS: Cure-Rite® 18 (purified), purity: 97.5%

Remark: Signed QA assurance statement provided

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Flag: Critical study for SIDS endpoint

12-OCT-2001 (8) (9)

- 18/28 -

Type: Cytogenetic assay

System of

testing: Chinese Hamster Ovary (CHO) Cells

Concentration: 2.500 to 20.000 ug/ml

Cytotoxic Conc.: concentration used based on mouse lymphoma L5178Y cells

Metabolic

activation: with and without

Result: positive

Method: other: according to Hinderer, R.K., B. Myhr, D.R. Jagannath,

S.M. Galloway, S.W. Mann, J.C.Riddle, and D.J. Brusick (1983)

Environ. Mutagen. 5:193-215

Year: 1983 GLP:

Test substance: other TS: Cure-Rite® 18 (purified); purity = 97.5%.

Remark: Signed QA assurance statement provided

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Flag: Critical study for SIDS endpoint

12-OCT-2001 (8) (10)

Type: DNA damage and repair assay

System of

testing: Escherichia coli strains W3110 (pol A+) and W3078 (pol A-)

Concentration: 100 to 5,000 ug/plate

Cytotoxic Conc.: With metabolic activation: 0.5 to 100 ug/plate (little or no

toxicity) Without metabolic activation: 0.5 to 100 ug/plate

(little or no toxicity)

Metabolic

activation: with and without

Result: positive

Method: other: according to Hinderer, R.K., B. Myhr, D.R. Jagannath,

S.M. Galloway, S.W. Mann, J.C.Riddle, and D.J. Brusick (1983)

Environ. Mutagen. 5:193-215

Year: 1983 GLP:

Test substance: other TS: Cure-Rite® 18.(purified); purity: 97.5%

Remark: Signed QA assurance statement provided

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Flag: Critical study for SIDS endpoint

12-OCT-2001 (8) (9)

Type: Mammalian cell gene mutation assay

System of

testing: BALB 3T3 Mouse Cells Concentration: 0.01000 to 0.20000 ug/ml

Cytotoxic Conc.: 0.488 ug/ml

Metabolic

activation: without Result: positive

Method: other: according to Hinderer, R.K., B. Myhr, D.R. Jagannath,

S.M. Galloway, S.W. Mann, J.C.Riddle, and D.J. Brusick (1983)

Environ. Mutagen. 5:193-215

Year: 1983 GLP:

Test substance: other TS: Cure-Rite® 18 (purified); purity = 97.5%.

- 19/28 -

Remark: Precipitation conc: >250 ug/ml Signed QA assurance statement provided.

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Flag: Critical study for SIDS endpoint

12-OCT-2001 (8) (11)

Type: Mouse lymphoma assay

System of

testing: Mouse Lymphoma cell line L5178Y TK+/-

Concentration: 1.250 to 25.0 ug/ml

Cytotoxic Conc.: With metabolic activation: Percent relative growth was 64.9%

at 1.560 ug/ml and 5.2% at 25.0 ug/ml

Without metabolic activation: Percent relative growth was

29.7% at 1.250 ug/ml and 7.7-11.2% at 5.0-20.0 ug/ml

Metabolic

activation: with and without

Result: positive

Method: other: according to Hinderer, R.K., B. Myhr, D.R. Jagannath,

S.M. Galloway, S.W. Mann, J.C.Riddle, and D.J. Brusick (1983)

Environ. Mutagen. 5:193-215

Year: 1983 GLP:

Test substance: other TS: Cure-Rite® 18 (purified); purity = 97.5%.

Remark: Precipitation conc: >250 ug/ml

Signed QA assurance statement provided.

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Flag: Critical study for SIDS endpoint

12-OCT-2001 (8) (9)

Type: Bacterial reverse mutation assay

System of

testing: Salmonella typhimurium strains TA-1535, TA-1537, TA-1538,

TA-98, TA-100

Concentration: 0.5 to 5,000 ug/plate

Cytotoxic Conc.: With metabolic activation: 0.5 to 100 ug/plate (little to no

toxicity)

Without metabolic activation: 0.5 to 100 ug/plate (little to

no toxicity)

Metabolic

activation: with and without

Result: negative

Method: other: according to Ames et al (1975) Mutation Res.

31:347-364; McCann et al. (1975) Proc. Nat. Acad. Sci.

72:5135-5139

Year: 1975 GLP:

Test substance: other TS: Commercial Cure-Rite® 18, purity: 95.6%

Remark: Signed QA assurance statement provided

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

12-OCT-2001 (8) (9)

- 20/28 -

Type: Bacterial reverse mutation assay

System of

testing: Salmonella typhimurium strains TA-1535, TA-1537, TA-1538,

TA-98, TA-100 and Saccharomyces strain D4

Concentration: 0.5 to 1,000 ug/plate

Cytotoxic Conc.: With metabolic activation: 1000 ug/plate (TA-98); 1000

ug/plate, 500 ug/plate, and 100 ug/plate (D4)

Without metabolic activation: 1000 ug/plate (TA-98); 1000

ug/plate, 500 ug/plate, and 100 ug/plate (D4)

Metabolic

activation: with and without

Result: negative

Method: other: according to Ames et al (1975) Mutation Res.

31:347-364; McCann et al. (1975) Proc. Nat. Acad. Sci.

72:5135-5139

Year: 1975 GLP:

Test substance: other TS: Commercial Cure-Rite® 18; purity: not noted

Remark: QA assurance statement provided.

Saccharomyces strain D4 not reported in referenced

publication.

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

12-OCT-2001 (8) (9)

Type: Bacterial reverse mutation assay

System of

testing: Escherichia coli strain WP2urvA

Concentration: 0.5 to 1,000 ug/plate

Cytotoxic Conc.: With metabolic activation: 0.5 to 100 ug/plate (little or no

toxicity) Without metabolic activation: 0.5 to 100 ug/plate

(little or no toxicity)

Metabolic

activation: with and without

Result: negative

Method: other: according to Hinderer, R.K., B. Myhr, D.R. Jagannath,

S.M. Galloway, S.W. Mann, J.C.Riddle, and D.J. Brusick (1983)

Environ. Mutagen. 5:193-215

Year: 1983 GLP:

Test substance: other TS: Commercial Cure-Rite® 18, purity: 95.6%

Remark: Signed QA assurance statement provided

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

12-OCT-2001 (8) (9)

- 21/28 -

Type: Cytogenetic assay

System of

testing: Chinese Hamster Ovary (CHO) Cells

Concentration: 0.313 to 5.000 ug/ml

Cytotoxic Conc.: concentration used based on mouse lymphoma L5178Y cells

Metabolic

activation: with and without

Result:

Method: other: according to Hinderer, R.K., B. Myhr, D.R. Jagannath,

S.M. Galloway, S.W. Mann, J.C.Riddle, and D.J. Brusick (1983)

Environ. Mutagen. 5:193-215

Year: 1983 GLP:

Test substance: other TS: Commercial Cure-Rite® 18; purity = 95.6%.

Remark: Signed QA assurance statement provided

Result: Genotoxic effects: With metabolic activation: negative

Without metabolic activation: positive

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

12-OCT-2001 (8) (10)

Type: DNA damage and repair assay

System of

testing: Escherichia coli strains W3110 (pol A+) and W3078 (pol A-)

Concentration: 100 to 5,000 ug/plate

Cytotoxic Conc.: With metabolic activation: 0.5 to 100 ug/plate (little or no

toxicity) Without metabolic activation: 0.5 to 100 ug/plate

(little or no toxicity)

Metabolic

activation: with and without

Result: positive

Method: other: according to Hinderer, R.K., B. Myhr, D.R. Jagannath,

S.M. Galloway, S.W. Mann, J.C.Riddle, and D.J. Brusick (1983)

Environ. Mutagen. 5:193-215

Year: 1983 GLP:

Test substance: other TS: Commercial Cure-Rite® 18, purity: 95.6%

Remark: Signed QA assurance statement provided

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

12-OCT-2001 (8) (9)

Type: Mammalian cell gene mutation assay

System of

testing: BALB 3T3 Mouse Cells Concentration: 0.05000 to 0.10000 ug/ml

Cytotoxic Conc.: 0.488 ug/ml

Metabolic

activation: without Result: negative

Method: other: according to Hinderer, R.K., B. Myhr, D.R. Jagannath,

S.M. Galloway, S.W. Mann, J.C.Riddle, and D.J. Brusick (1983)

Environ. Mutagen. 5:193-215

Year: 1983 GLP:

Test substance: other TS: Commercial Cure-Rite® 18; purity = 95.6%.

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Remark: Signed QA assurance statement provided

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

12-OCT-2001 (8) (11)

Type: Mammalian cell gene mutation assay

System of

testing: BALB 3T3 Mouse Cells Concentration: 0.00625 to 0.10000 ug/ml

Cytotoxic Conc.: 0.244 ug/ml

Metabolic

activation: without Result: positive

Method: other: according to Hinderer, R.K., B. Myhr, D.R. Jagannath,

S.M. Galloway, S.W. Mann, J.C.Riddle, and D.J. Brusick (1983)

Environ. Mutagen. 5:193-215

Year: 1983 GLP:

Test substance: other TS: Commercial Cure-Rite® 18, purity: Not noted

Remark:

Weakly active.

Signed QA assurance statement provided. Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

12-OCT-2001 (8) (11)

Type: Mouse lymphoma assay

System of

testing: Mouse Lymphoma cell line L5178Y TK+/-

Concentration: 0.313 to 35.0 ug/ml

Cytotoxic Conc.: With metabolic activation: Percent relative growth was 78.1%

at 20.0 ug/ml and 5.7% at 35.0 ug/ml

Without metabolic activation: Percent relative growth was 25.6%

at 0.313 ug/ml and 3.8% at 1.880 ug/ml

Metabolic

activation: with and without

Result: positive

Method: other: according to Hinderer, R.K., B. Myhr, D.R. Jagannath,

S.M. Galloway, S.W. Mann, J.C.Riddle, and D.J. Brusick (1983)

Environ. Mutagen. 5:193-215

Year: 1983 GLP:

Test substance: other TS: Commercial Cure-Rite® 18; purity = 95.6%.

Remark:

Weakly active.

Signed QA assurance statement provided.

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

12-OCT-2001 (8) (9)

- 23/28 -

Type: Mouse lymphoma assay

System of

testing: Mouse Lymphoma cell line L5178Y TK+/-

Concentration: 0.313 to 50.0 ug/ml

Cytotoxic Conc.: With metabolic activation: Percent relative growth was 43.2%

at 12.50 ug/ml and 4.2% at 50.0 ug/ml

Without metabolic activation: Percent relative growth was

80.9% at 0.313 ug/ml and 7.9% at 1.880 ug/ml

Metabolic

activation: with and without

Result:

Method: other: according to Hinderer, R.K., B. Myhr, D.R. Jagannath,

S.M. Galloway, S.W. Mann, J.C.Riddle, and D.J. Brusick (1983)

Environ. Mutagen. 5:193-215

Year: 1983 GLP:

Test substance: other TS: Commercial Cure-Rite® 18; purity: Not noted

Remark: Precipitation conc: 1250 ug/ml Signed QA assurance statement provided.

Result: With metabolic activation: weakly active Without metabolic activation: negative

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

12-OCT-2001 (8) (9)

5.6 Genetic Toxicity 'in Vivo'

Type: Dominant lethal assay

Species: rat Sex: male/female

Strain: Sprague-Dawley

Route of admin.: gavage

Exposure period: 56 consecutive days to males

Doses: 0, 6.25, 12.5, or 25 mg/kg. (0.25 mg/kg triethylenemelamine

positive control)

Result: negative

Method: other: according to Hinderer, R.K., M. Knickerbocker, and F.J.

Koschier (1982) Toxicol. Appl. Pharmacol. 62:335-341.

Year: 1982 GLP: yes

Test substance: other TS: Commercial Cure-Rite® 18; Purity = 95.6%

Result: A significant depression in body weight gain was observed in

the males administered the highest dose. Similar pregnancy rates were observed in all test groups compared with the controls. No evidence of dominant lethal mutations were observed in the test groups. In the TEM positive controls, the number of implantation sites and preimplantation loses were significantly decreased, and the number of early fetal deaths per pregnant female were significantly elevated.

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Flag: Critical study for SIDS endpoint

12-OCT-2001 (12)

- 24/28 -

5.7 Carcinogenicity

Species: Sex: male/female

Strain: Sprague-Dawley Route of admin.: oral feed Exposure period: 2 years

Frequency of

treatment: continuous daily

Post. obs.

period: none

0, 20, 60, 200, or 600 ppm Doses:

Result:

Control Group: yes, concurrent no treatment

Method: other: according to Hinderer, R.K., G.R. Lankas, A.L.

Knezevich, and C.S. Auletta (1986). Toxicol. Appl. Pharmacol.

82:521-531

Year: 1986 GLP:

Test substance: other TS: Commercial Cure-Rite® 18; purity = 96.8%

Result: A compound related increase in urothelial tumors, kidney

weights, non-neoplastic urinary tract abnormalities, and rales was observed in the high dose males and females. Body weights

also were significantly lower in the high dose males and

females. No compound-related effects on hematology, clinical

chemistry, or urinalysis were noted.

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Flag: Critical study for SIDS endpoint

12-OCT-2001 (7)

5.8 Toxicity to Reproduction

Type: Fertility

Species: Sex: male/female

Strain: Sprague-Dawley Route of admin.: oral feed Exposure Period: 12 weeks

Frequency of

treatment: daily Premating Exposure Period

Males were sacrificed over a 6-7 day period following the male:

21-day mating period.

Duration of test: 12 weeks

0, 60, 200, or 600 ppm. Doses: Control Group: yes, concurrent no treatment

NOAEL Parental: 200 ppm NOAEL F1 Offspr.: 600 ppm

Method: other: according to Hinderer, R.K., B.Y. Cockrell, S.M.

Debanne, and P.T. Goad. (1987). Fund. Appl. Toxicol.

9:763-772.

Year: 1987 GLP: yes

Test substance: other TS: Commercial Cure-Rite® 18; purity = 98.0%.

Result: NOEL Parental: 200 ppm based on a slight, but generally not

statistically significant body weight reduction at 600 ppm

- 25/28 -

No evidence of a compound-related effect on mating, fertility, gestation length, number of implants or live births, pup

growth, or survival was observed.

No morphological changes in the testes from the high dose males was observed by either light or electron microscopy.

General parental toxicity: A slight but generally

nonstatistically significant decrease in body weights in the

test males.

Toxicity to offspring: None

Reliability: (1) valid without restriction

GLP study, Meets generally accepted scientific standards, well

documented and acceptable for assessment.

Flag: Critical study for SIDS endpoint

12-OCT-2001 (13)

5.9 Developmental Toxicity/Teratogenicity

See IUCLID data sets on CAS# 95-31-8; 102-77-2; 95-33-0; 4979-32-2

5.10 Other Relevant Information

-

5.11 Experience with Human Exposure

-

- 26/28 -

Date: 12-OCT-2001
6. References ID: 13752-51-7

1. 1.01.01.01.01

(1) Meylan W. and Howard P. (1999) EPIWin Modeling Program. Syracuse Research Corporation. Environmental Science Center, 6225 Running Ridge Road, North Syracuse, NY 13212-2510.

- (2) BFGoodrich MSDS
- (3) Bio/dynamics Inc., East Millstone, NJ (1980) Project #6216-80
- (4) Bio/dynamics Inc., East Millstone, NJ (1980)
- (5) Hill Top Research, Inc., Miamiville, Ohio (1971)
- (6) Biosearch Inc, Philadelphia, PA(1981)
- (7) Hinderer, R.K., G.R. Lankas, A.L. Knezevich, and C.S. Auletta (1986). Toxicol. Appl. Pharmacol. 82:521-531
- (8) Hinderer, R.K., B. Myhr, D.R. Jagannath, S.M. Galloway, S.W. Mann, J.C.Riddle, and D.J. Brusick (1983) Environ. Mutagen. 5:193-215
- (9) Litton Bionetics, Inc Report Project Number 20988 (1979)
- (10) Litton Bionetics, Inc Report Project Number 20990 (1979)
- (11) Litton Bionetics, Inc Report Project Number 20992 (1980)
- (12) Hinderer, R.K., M. Knickerbocker, and F.J. Koschier (1982). Toxicol. Appl. Pharmacol. 62:335-341.
- (13) Hinderer, R.K., B.Y. Cockrell, S.M. Debanne, and P.T. Goad. (1987). Fund. Appl. Toxicol. 9:763-772.

- 27/28 -

7. Risk Assessment Date: 12-OCT-2001 ID: 13752-51-7

7.1 End Point Summary

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7.2 Hazard Summary

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7.3 Risk Assessment

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- 28/28 -

IUCLID

Data Set

Existing Chemical ID: 4979-32-2 CAS No. 4979-32-2

EINECS Name N,N-dicyclohexylbenzothiazole-2-sulphenamide

EINECS No. 225-625-8

TSCA Name 2-Benzothiazolesulfenamide, N,N-dicyclohexyl-

Molecular Formula C19H26N2S2

Producer Related Part

Company:

Creation date: 26-APR-2001

Substance Related Part

Company:

Creation date: 26-APR-2001

Memo: Data for RAPA Sulfenamide Accelerators category

Printing date: 18-OCT-2001

Revision date:

Date of last Update: 18-OCT-2001

Number of Pages: 28

Chapter (profile): Chapter: 1, 2, 3, 4, 5, 7

Reliability (profile): Reliability: without reliability, 1, 2, 3, 4

Flags (profile): Flags: without flag, confidential, non confidential, WGK

(DE), TA-Luft (DE), Material Safety Dataset, Risk

Assessment, Directive 67/548/EEC, SIDS

Date: 18-OCT-2001

1. General Information ID: 4979-32-2

 $1.0.1\ \text{OECD}$ and Company Information

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1.0.2 Location of Production Site

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1.0.3 Identity of Recipients

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1.1 General Substance Information

Substance type: organic Physical status: solid

Purity: >= 95 % w/w

Remark: cooperating companies:

Bayer Antwerpen N.V., Belgium AKZO Chemicals, Netherlands Monsanto Europe N.V., Belgium

26-APR-2001

1.1.0 Details on Template

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1.1.1 Spectra

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1.2 Synonyms

 $\begin{tabular}{ll} N, N-dicyclohexyl-2-benzothiazolesulfenamide \\ 26-APR-2001 \end{tabular}$

Sanotcure DCBS

Source: Bayer AG Leverkusen

07-JUN-1994

Vulkacit DZ

Source: Bayer AG Leverkusen

07-JUN-1994

1.3 Impurities

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- 1/28 -

Date: 18-OCT-2001 ID: 4979-32-2 1. General Information

1.4 Additives

CAS-No: EINECS-No: EINECS-Name:

Remark: Additives may be contained in the substance as marketed in

order to reduce dust formation during use.

26-APR-2001

1.5 Quantity

1.6.1 Labelling

1.6.2 Classification

1.7 Use Pattern

Type: type

Category: Use resulting in inclusion into or onto matrix

26-APR-2001

Type: industrial

Category: Polymers industry Bayer AG Leverkusen Source:

26-MAY-1994

Type: use

Category: Vulcanizing agents Bayer AG Leverkusen Source:

26-MAY-1994

1.7.1 Technology Production/Use

1.8 Occupational Exposure Limit Values

1.9 Source of Exposure

1.10.1 Recommendations/Precautionary Measures

- 2/28 -

Date: 18-OCT-2001

1. General Information ID: 4979-32-2

1.10.2 Emergency Measures

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1.11 Packaging

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1.12 Possib. of Rendering Subst. Harmless

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1.13 Statements Concerning Waste

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1.14.1 Water Pollution

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1.14.2 Major Accident Hazards

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1.14.3 Air Pollution

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1.15 Additional Remarks

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1.16 Last Literature Search

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1.17 Reviews

-

1.18 Listings e.g. Chemical Inventories

-

- 3/28 -

2.1 Melting Point

Value: >= 96 degree C

Decomposition: no
Sublimation: no
Method: other
Year: 1982
GLP: no

Remark: sample of technical purity

Source: Bayer AG Leverkusen

30-APR-1992 (1)

Value: = 103.5 degree C

Decomposition: no
Sublimation: no
Method: other
Year: 1978
GLP: no

Remark: high purity sample Source: Bayer AG Leverkusen

30-APR-1992 (1)

2.2 Boiling Point

Value: >= 200 degree C at 1013 hPa

Decomposition: yes
Method: other
Year: 1976
GLP: no

Remark: DCBS was heated in an open glass vessel under air and

normalpressure with a heating rate of 10 degree C per min, and in a closed glass vessel with a heating rate of 5 degree C per min. In both experiments a very strong exothermic decomposition could be observed at temperatures of >= 200

degree C using DTA.

Thermogravimetric analysis showed that when a DCBS sample was kept at a constant temperature of 230 degree C on a balance, after 5 min more than 50 % w/w of the sample were volatilized. After 30 and 45 min approx. 20 % w/w of the

sample were still on the balance.

Source: Bayer AG Leverkusen

04-MAY-1992 (1)

2.3 Density

Type: bulk density

Value: ca. 1.2 g/cm3 at 20 degree C

Method: other Year: 1982 GLP: no

Source: Bayer AG Leverkusen

30-APR-1992 (1)

- 4/28 -

2. Physico-chemical Data

2.3.1 Granulometry

2.4 Vapour Pressure

ca. .00075 hPa at 120 degree ${\tt C}$ Value:

Method: other (measured): comparable to OECD Guide-line 104

Year: 1978 GLP: no

Bayer AG Leverkusen Source:

30-APR-1992 (1)

Value: ca. .0035 hPa at 140 degree C

Method: other (measured): comparable to OECD Guide-line 104

Year: 1978 GLP: no

Source: Bayer AG Leverkusen

30-APR-1992 (1)

2.5 Partition Coefficient

log Pow: 5.951 at 25 degree C

Method: other (calculated): KOWWIN Program (v1.65)

Year: 1999 GLP: no

Testsubstance: other TS: molecular structure Reliability: (2) valid with restrictions Accepted calculation method

18-OCT-2001 (2)

log Pow: 4.8

Method: OECD Guide-line 107 "Partition Coefficient (n-octanol/water),

Flask-shaking Method"

Year:

Source: Bayer AG Leverkusen

04-DEC-1995 (3)

2.6.1 Water Solubility

Value: ca. 30 mg/l at 25 degree C of very low solubility Qualitative:

Method: other Year: 1988 GLP: no

Source: Bayer AG Leverkusen

30-APR-1992 (1)

2.6.2 Surface Tension

- 5/28 -

Date: 18-OCT-2001
2. Physico-chemical Data

Date: 18-OCT-2001

1D: 4979-32-2

2.7 Flash Point

Value: ca. 180 degree C

Type: closed cup

Method: other: DIN 51758

Year: 1989 GLP: no

Remark: The substance being solid, flash point determination

according to the EEC-Directive cannot be carried out. The determination was done using a molten sample following

method DIN 51758.

Source: Bayer AG Leverkusen

04-MAY-1992 (1)

2.8 Auto Flammability

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2.9 Flammability

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2.10 Explosive Properties

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2.11 Oxidizing Properties

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2.12 Additional Remarks

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- 6/28 -

3. Environmental Fate and Pathways

3.1.1 Photodegradation

Type: air INDIRECT PHOTOLYSIS Sensitizer: OH

Conc. of sens.: 1560000 molecule/cm3

Rate constant: .000000001138512 cm3/(molecule * sec)

Degradation: 50 % after 1.1 hour(s)

Method: other (calculated): AOP Program (v1.89) 1999 Year:

Test substance: other TS: molecular structure Reliability: (2) valid with restrictions Accepted calculation method

18-OCT-2001 (2)

3.1.2 Stability in Water

Type: abiotic Method: other

Year: 1988 GLP: no

Test substance: as prescribed by 1.1 - 1.4

Result: DCBS hydrolyzed slowly at a temperature of 100 degree C,

2-mercaptobenzothiazole (MBT), MBT-sulfonic acid,

2-hydroxybenzothiazole and dicyclohexylamine were identified as decomposition products. Addition of strong bases or acids

accelerated the reaction.

Source: Bayer AG Leverkusen

30-APR-1992 (1)

3.1.3 Stability in Soil

3.2 Monitoring Data (Environment)

3.3.1 Transport between Environmental Compartments

fugacity model level III Type:

Media: other: air - water - soil - sediment

Air (Level I): Water (Level I): Soil (Level I): Biota (L.II/III): Soil (L.II/III):

Method: other: Level III Fugacity Model

Year: 1999

Result: Media Distribution Half-Life Emissions Fugacity (percent) (hr) (kg/hr) (atm) Air 0.0457 2.25 1000 2.42e-013 Water 7.15 900 1000 7.39e-015

Soil 39.9 900 1000 8.32e-017

Date: 18-OCT-2001 3. Environmental Fate and Pathways ID: 4979-32-2

Sediment 52.9 3.6e+003 0 4.97e-015

> Persistence Time: 1.45e+003 hr Reaction Time: 1.65e+003 hr Advection Time: 1.15e+004 hr Percent Reacted: 87.5

Percent Advected: 12.5

(2) valid with restrictions Reliability:

Accepted calculation method

18-OCT-2001 (2)

3.3.2 Distribution

3.4 Mode of Degradation in Actual Use

3.5 Biodegradation

Type: aerobic

Inoculum: predominantly domestic sewage
Concentration: 100 mg/l related to Test substance
Degradation: ca. 2 % after 28 day

Result: under test conditions no biodegradation observed

Directive 84/449/EEC, C.7 "Biotic degradation - modified MITI Method:

test"

Year: 1989 GLP: no

Test substance: other TS: commercial product Source: Bayer AG Leverkusen

04-DEC-1995 (1)

Type: aerobic

Inoculum: other: Japanese standard activated sludge

Concentration: 100 mg/l related to Test substance
Degradation: 0 % after 28 day

Method: OECD Guide-line 301 C "Ready Biodegradability: Modified MITI

Test (I)"

1994 Year: GLP: yes

Test substance:

Source: Bayer AG Leverkusen

04-DEC-1995 (4)

3.6 BOD5, COD or BOD5/COD Ratio

Remark: ThOD: 2300 mg/g Source: Bayer AG Leverkusen

09-DEC-1993 (1)

- 8/28 -

Date: 18-OCT-2001
3. Environmental Fate and Pathways ID: 4979-32-2

3.7 Bioaccumulation

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3.8 Additional Remarks

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- 9/28 -

AQUATIC ORGANISMS

4.1 Acute/Prolonged Toxicity to Fish

semistatic Type:

Species: Oryzias latipes (Fish, fresh water)

Exposure period: 96 hour(s)

mg/lAnalytical monitoring: no

LC50: > 1000

Method: other: OECD Guideline 203

1981 GLP: no Year:

Test substance: other TS: 99.9 %

Remark: Stock solution was prepared with ethanol with

ultrasonication.

Source: Bayer AG Leverkusen

Reliability: (1) valid without restriction

Guideline study

18-OCT-2001 (3)(5)

Type: static

Brachydanio rerio (Fish, fresh water) Species:

Exposure period: 96 hour(s)

Analytical monitoring: no Unit: mq/1

NOEC: 15 Method: other

Year: 1988 GLP: no

Test substance: other TS: commercial product

Remark: The substance was dispersed in water equivalent to a

> concentration of 1 q/l, stirred for 2 h and subsequently filtered. The filtrate was tested on Brachydanio rerio: no observed effects in the undiluted sample (DOC of the

filtrate : 15 mg/l).

Bayer AG Leverkusen Source:

26-JAN-1995 (1)

4.2 Acute Toxicity to Aquatic Invertebrates

Type:

Daphnia magna (Crustacea) Species:

Exposure period: 24 hour(s)

Analytical monitoring: no Unit: mg/1

EC50: > 1000

Method: other: OECD Guideline 202

1984 GLP: no Year:

other TS: 99.9 % Test substance:

Remark: static

stock solution was prepared with DMSO:HCO-40 = 9:1

Bayer AG Leverkusen Source:

(1) valid without restriction Reliability:

Guideline study

18-OCT-2001 (3)

- 10/28 -

4.3 Toxicity to Aquatic Plants e.g. Algae

Species: Selenastrum capricornutum (Algae)

Endpoint: biomass
Exposure period: 72 hour(s)

Unit: mg/l Analytical monitoring: no

NOEC: 10 EC50: 16

Method: other: OECD Guideline 201

Year: 1984 GLP: no

Test substance: other TS: 99.9 %

Remark: Stock solution was prepared with DMSO:HCO-40 = 9:1

Source: Bayer AG Leverkusen

Reliability: (1) valid without restriction

Guideline study

18-OCT-2001 (3) (5)

4.4 Toxicity to Microorganisms e.g. Bacteria

Type: aquatic

Species: activated sludge

Exposure period: 3 hour(s)

Unit: mg/l Analytical monitoring: no

EC0: >= 10000

Method: ISO 8192 "Test for inhibition of oxygen consumption by

activated sludge"

Year: 1988 GLP: no

Test substance: other TS: commercial product

Remark: The substance was dispersed in the test medium equivalent to

a concentration of 10 g/l.

Source: Bayer AG Leverkusen

09-DEC-1993 (1)

- 11/28 -

- 4.5 Chronic Toxicity to Aquatic Organisms
- 4.5.1 Chronic Toxicity to Fish

-

4.5.2 Chronic Toxicity to Aquatic Invertebrates

Species: Daphnia magna (Crustacea)

Endpoint: reproduction rate

Exposure period: 21 day

Unit: mg/l Analytical monitoring: no

NOEC: 10 LOEC: 18 EC50: 40

Method: other: OECD Guideline 202 (1984)

Year: 1994 GLP: no

Test substance: other TS: 99.9 %

Remark: 21d-EC50 (immobility): 140 mg/l

static

Stock solution was prepared with DMSO:HCO-40 = 9:1

Source: Bayer AG Leverkusen

31-MAY-1996 (3) (5)

TERRESTRIAL ORGANISMS

4.6.1 Toxicity to Soil Dwelling Organisms

-

4.6.2 Toxicity to Terrestrial Plants

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4.6.3 Toxicity to other Non-Mamm. Terrestrial Species

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4.7 Biological Effects Monitoring

-

4.8 Biotransformation and Kinetics

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4.9 Additional Remarks

-

- 12/28 -

5.1 Acute Toxicity

5.1.1 Acute Oral Toxicity

Type: LD50 Species: rat

Strain:

Sex: male/female

Number of
Animals:
Vehicle:
Value:

Method: OECD Guide-line 401 "Acute Oral Toxicity" Year: GLP: yes

Test substance: other TS: purity 99.2 %

Remark: Fatal cases were found in males at the doses of more than

2367 mg/kg and in females of more than 1401 mg/kg. But, no

dose-related mortalities were observed.

value: > 1,821 mg/kg in male; > 1,077 mg/kg in female

Source: Bayer AG Leverkusen

Reliability: (1) valid without restriction

GLP Guideline study

18-OCT-2001 (6)

Type: LD50 Species: rat

Strain:
Sex:
Number of
Animals:
Vehicle:

Value: = 10000 mg/kg bw

Method:

Year: GLP:

Test substance:

Source: Bayer AG Leverkusen

20-OCT-1993 (7)

Type: LD50 Species: rat

Strain:
Sex:
Number of
 Animals:
Vehicle:

Value: = 6420 mg/kg bw

Method:

Year: GLP:

Test substance:

Source: Bayer AG Leverkusen

20-OCT-1993 (8)

- 13/28 -

Type: LD50 Species: rat

Strain:
Sex:
Number of
 Animals:
Vehicle:

Value: = 8500 mg/kg bw

Method:

Year: GLP:

Test substance:

Source: Bayer AG Leverkusen

10-MAY-1994 (9)

Type: LD50 Species: rat

Strain:
Sex:
Number of
Animals:
Vehicle:

Value: > 5000 mg/kg bw

Method:

Year: GLP:

Test substance:

Source: Monsanto

Bayer AG Leverkusen

06-JUN-1994 (10)

5.1.2 Acute Inhalation Toxicity

-

5.1.3 Acute Dermal Toxicity

Type: LD50 Species: rabbit

Strain:
Sex:
Number of
Animals:
Vehicle:

Value: > 2000 mg/kg bw

Method:

Year: GLP:

Test substance: other TS: purity = 96%

Source: Monsanto

Bayer AG Leverkusen

26-APR-2001 (10)

- 14/28 -

Date: 18-OCT-2001
5. Toxicity

Date: 18-OCT-2001

5.1.4 Acute Toxicity, other Routes

Type: LD50 Species: rat

Strain:
Sex:
Number of
 Animals:
Vehicle:

Route of admin.: s.c.

Value: > 5000 mg/kg bw

Method:

Year: GLP:

Test substance:

Source: Bayer AG Leverkusen

Test substance: DCBS of technical and analytical grade was tested.

20-OCT-1993 (11)

5.2 Corrosiveness and Irritation

5.2.1 Skin Irritation

Species: rabbit

Concentration:

Exposure:
Exposure Time:
Number of
Animals:
PDII:
Result:

EC classificat.:

Method:

Year: GLP:

Test substance:

Remark: 20 mg/24h

effect: moderate

Source: Bayer AG Leverkusen

20-OCT-1993 (12)

Species: rabbit

Concentration:

Exposure:
Exposure Time:
Number of
Animals:
PDII:
Result:

EC classificat.:

Method: other: exposure time: 24 h (no further data)

Year: GLP:

Test substance:

Remark: effect: practically non-irritating

- 15/28 -

Date: 18-OCT-2001
5. Toxicity

Date: 18-OCT-2001

Source: Bayer AG Leverkusen

26-APR-2001 (10)

5.2.2 Eye Irritation

Species: rabbit

Concentration:

Dose:

Exposure Time:
Comment:
Number of
Animals:
Result:

EC classificat.:

Method:

Year: GLP:

Test substance:

Remark: 500 mg/24 h

effect: mild

Source: Bayer AG Leverkusen

20-OCT-1993 (12)

Species: rabbit

Concentration:

Dose:

Exposure Time:
Comment:
Number of
Animals:

Result:

EC classificat.:

Method: other: exposure time: 24 h (no further data)

Year: GLP:

Test substance:

Remark: effect: practically non-irritating

Source: Monsanto

Bayer AG Leverkusen

06-JUN-1994 (10)

- 16/28 -

5.3 Sensitization

Type: Guinea pig maximization test

Species: guinea pig

Number of
Animals:
Vehicle:

Result: not sensitizing

Classification:

Method:

Year: GLP:

Test substance: other TS: Santocure DCBS; purity = 98%

Source: Monsanto

Bayer AG Leverkusen

26-APR-2001 (13)

Type: Patch-Test Species: human

Number of
Animals:
Vehicle:
Result:

Classification:

Method:

Year: GLP:

Test substance:

Remark: 2/135 showed an allergic reaction towards 1 %

Cyclohexylbenzothiazyl-sulfenamid

Source: Bayer AG Leverkusen

30-JUL-1998 (14)

5.4 Repeated Dose Toxicity

Species: rat Sex: male/female

Strain: other: Crj: CD (SD)
Route of admin.: other: oral gavage

Exposure period: males: 44 days including 14 days before mating; females: from

14 days before mating to day 3 of lactation

Frequency of

treatment: 7 days/week

Post. obs. period:

Doses: 0, 6, 25, 100, 400 mg/kg (10 animals/group)

Control Group: yes, concurrent vehicle

Method: other: OECD Combined Repeat dose and

reproductive/Developmental Screening Toxicity Test

Year: GLP: yes

Test substance: other TS: purity 99.2 %

Result: In the cage side observation, salivation were noted in the

400 mg/kg male group, and decreased locomotor activity were noted in 100 mg/kg or more female groups. At a dose of 400 mg/kg, food consumption decreased prior to the mating and during the pregnancy, and body weight gain decreased at late stage of pregnancy. In urinalysis, increased ketone bodies

- 17/28 -

were noted in 400 mg/kg males. There are no hematological changes between the treated and control groups of both sexes. Increased relative kidney weights and decreased absolute thymus weights were noted in 400 mg/kg male group and 400 mg/kg both male and female groups, respectively. In histopathological examination, fatty degeneration of the renal tubular epithelia, vacuolation of the adreno-cortical cells and atrophy of the spleen were observed at doses of 100 and 400 mg/kg female groups. Fatty degeneration of liver cells were observed in 400 mg/kg male group, and congestion of liver was noted in 400 mg/kg female group. Also, hyaline droplets in the renal tubular epithelia were observed in 100

and 400 mg/kg male groups.

Source: Bayer AG Leverkusen

Reliability: (1) valid without restriction

GLP Guideline study

18-OCT-2001 (15)

Species: rat Sex: male/female

Strain: Sprague-Dawley

Route of admin.: oral feed

Exposure period:

Frequency of

treatment: daily

Post. obs.

period: no data

Doses: 2000, 3000, 5000, 7500 or 10000 ppm (ca. 133, 200, 333, 500 or

667 mg/kg bw/d)

Control Group: yes

Method:

Year: GLP:

other TS: Santocure DCBS Test substance:

Result: no significant changes related to treatment were found

> in hematology or clinical chemistry evaluations, terminal organ weights, or organ/body weight ratios or gross necropsy examinations; dose-related depression in body weight gain and reduced feed consumption on a mg/kg bw/d basis was noted in all treatment groups in

comparison to controls

Source: Monsanto

Bayer AG Leverkusen

18-OCT-2001 (16)

- 18/28 -

Species: rat Sex: male/female

Strain: Sprague-Dawley
Route of admin.: oral feed
Exposure period: 3 months

Frequency of

treatment: daily

Post. obs.

period: no data

Doses: 2500 or 5000 ppm (ca. 167 or 333 mg/kg bw/d)

Control Group: yes

Method:

Year: GLP: yes
Test substance: other TS: Santocure DCBS; purity = 96%
Remark: species: unspecified, probably rat

Result: reduced body weight gain and reduced food consumption in

both sexes in both treatment groups were observed; no target organ toxicity or histopathological findings were sug-

gested

Source: Monsanto

Bayer AG Leverkusen

18-OCT-2001 (17)

Species: rat Sex: male

Strain: no data Route of admin.: inhalation

Exposure period: 15 d

Frequency of

treatment: 2h/d

Post. obs.

period: no data
Doses: 340-400 mg/m3
Control Group: other: no data

Method:

Year: GLP:

Test substance:

Result: No effect except mucous membrane irritaion were observed. No

pronounced liver or kidney changes were observed.

Source: Bayer AG Leverkusen

18-OCT-2001 (9)

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5.5 Genetic Toxicity 'in Vitro'

Type: Bacterial reverse mutation assay

System of

testing: S. typhimurium TA 98, TA 100, TA 1535, TA 1537

Concentration: 0, 312.5, 625, 1250, 2500, 5000 ug/plate

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method: other: Japanese Guideline for Screening Mutagenicity testing

of chemicals

Year: GLP: yes
Test substance: other TS: commercial, purity: 99.5 %

Remark: procedure: plate incorporation method

Result: cytotoxicity conc: with and without metabolic activation:

5000 ug/plate; precipitation conc: 312.5 ug/plate

Source: Bayer AG Leverkusen

Reliability: (1) valid without restriction
Meets National standards method

18-OCT-2001 (18)

Type: Bacterial reverse mutation assay

System of

testing: E. coli WP 2 uvrA

Concentration: 0, 312.5, 625, 1250, 2500, 5000 ug/plate

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method: other: Japanese Guideline for Screening Mutagenicity testing

of chemicals

Year: GLP: yes
Test substance: other TS: commercial, purity: 99.5 %
Remark: procedure: plate incorporation method

Result: cytotoxicity conc: with and without metabolic activation:

5000 ug/plate, precipitation conc: 312.5 ug/plate

Source: Bayer AG Leverkusen

Reliability: (1) valid without restriction
Meets National standards method

18-OCT-2001 (18)

Type: Cytogenetic assay

System of testing:

Concentration: -S9 (continuous treatment) 0, 0.21, 0.41, 0.82 mg/ml; -S9

(short-term treatment) 0, 0.9, 1.8, 3.5 mg/ml; +S9 (short-term

treatment) 0, 0.9, 1.8, 3.5 mg/ml

Cytotoxic Conc.:

Metabolic

activation: with and without

Result:

Method: other: Japanese Guideline for Screening Mutagenicity testing

of chemicals

Year: GLP: yes

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Test substance: other TS: commerical, purity: 99.5 %

Result: cytoxtoxicity conc: with and without metabolic activation: >

3.5 mg/ml; genotoxic effects: clastogenicity with and

without metabolic activation: negative, polyploid induction:

with metabolic activation: negative, without metabolic

activation: positive

Source: Bayer AG Leverkusen

Reliability: (1) valid without restriction

Meets National standards method

18-OCT-2001 (18)

Type: Ames test

System of

testing: S. typhimurium TA 100, TA 98

Concentration:
Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method:

Year: GLP:

Test substance:

Source: Bayer AG Leverkusen

04-MAY-1992 (19)

Type: Ames test

System of

testing: Salmonella typhimurium (no further data)

Concentration:
Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method:

Year: GLP: no data

Test substance:

Source: Monsanto

Bayer AG Leverkusen

26-APR-2001 (20)

Type: HGPRT assay

System of

testing: Chinese hamster ovary cells

Concentration: up to 500 ug/ml

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method:

Year: GLP:

Test substance: other TS: Santocure DCBS; purity = 96%

Source: Monsanto

Bayer AG Leverkusen

26-APR-2001 (21)

Date: 18-OCT-2001
5. Toxicity

Date: 18-OCT-2001

Type: Unscheduled DNA synthesis

System of

testing: primary rat hepatocytes
Concentration: up to and including 50 ug/ml

Cytotoxic Conc.:

Metabolic

activation:

Result: negative

Method:

Year: GLP:

Test substance: other TS: Santocure DCBS

Source: Monsanto

Bayer AG Leverkusen

06-JUN-1994 (22)

5.6 Genetic Toxicity 'in Vivo'

Type: Cytogenetic assay

Species: rat Sex: male/female

Strain: no data Route of admin.: gavage

Exposure period: single administration

Doses: 1000 mg/kg bw Result: negative

Method:

Year: GLP: yes
Test substance: other TS: Santocure DCBS; purity = 96%

Result: Santocure DCBS did not produce chromosome damage as mea-

sured by significant increases in chromosome aberrations or chromosome number as compared to concurrent controls

in the rat bone marrow assay

Source: Monsanto

Bayer AG Leverkusen

26-APR-2001 (23)

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5.7 Carcinogenicity

Species: rat Sex: male/female

Strain: Wistar Route of admin.: s.c. Exposure period: 413 d

Frequency of

treatment: once a week

Post. obs.

period: entire lifetime

Doses: 1000 mg/kg bw/week, 20000 mg/kg bw (total amount)

Result:

Control Group: other: yes Method: other

Year: 1975 GLP: no

Test substance: as prescribed by 1.1 - 1.4

Remark: 20 animals of each sex, technical and analytical grade DCBS

was administered.

Result: No signs of systemic toxicity were reported, there was no

difference between the survival of the control group and the dose group, an increased number of sarcomas located at the

injection site was observed in all dose groups.

Source: Bayer AG Leverkusen

30-JUL-1998 (11)

5.8 Toxicity to Reproduction

Type: other

Species: rat Sex: male/female

Strain: other: Crj: CD (SD)
Route of admin.: other: oral gavage

Exposure Period: males: 44 days including 14 days before mating, females: from

14 days before mating to day 3 of lactation

Frequency of

treatment: 7 days/week
Premating Exposure Period
male: 14 days
female: 14 days

Duration of test:

Doses: 0, 6, 25, 100 or 400 mg/kg (10 animals/sex/group)

Control Group: yes, concurrent vehicle

NOAEL Parental: 100 mg/kg bw NOAEL F1 Offspr.: 100 mg/kg bw

Method: other: OECD Combined Repeat dose and reproductive/Development

Screening Toxicity Test

Year: GLP: yes

Test substance: other TS: purity 99.2 %

Result: There were no effects indicative of toxicity on male

reproductive performance. Toxic effects were revealed in female and pups at doses of 400 mg/kg. There was decreased number of the corpus lutea in accompany of decreases in number of implantation sites and litter size. One dam died during the delivery and another two had prolonged gestation length. All dams lost their litters at delivery or by day 4

- 23/28 -

of lactation. Therefore, there were decreases in

reproduction/development parameters such as gestation index, number of live pups at birth, live birth index and viability index on day 4 of lactation. There were no effects on the

mating and fertility, and morphogenesis in pups.

Source: Bayer AG Leverkusen

Reliability: (1) valid without restriction

GLP Guideline study

18-OCT-2001 (15)

5.9 Developmental Toxicity/Teratogenicity

Species: rat Sex: male/female

Strain: other: Crj: CD (SD)

Route of admin.: gavage

Exposure period: males: 44 days including 14 days before mating, females: from

14 days before mating to day 3 of lactation

Frequency of

treatment: 7 days/week

Duration of test:

Doses: 0, 6, 25, 100 or 400 mg/kg (10 animals/sex/group)

Control Group: yes, concurrent vehicle

Method: other: OECD Combined Repeat dose and reproductive/Development

Screening Toxicity Test

Year: GLP: yes

Test substance: other TS: purity 99.2 %

Result: Toxic effects were revealed in female and pups at doses of 400

mg/kg. There were no effects on the mating and fertility, and

morphogenesis in pups.

One dam died during the delivery and another two had prolonged gestation length. All dams lost their litters at delivery or by day 4 of lactation. Therefore, there were decreases in reproduction/development parameters such as gestation index, number of live pups at birth, live birth index and viability

index on day 4 of lactation.

Reliability: (1) valid without restriction

GLP Guideline study

18-OCT-2001 (15)

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Date: 18-OCT-2001
5. Toxicity

Date: 18-OCT-2001

Species: other: chicken Sex:

Strain: no data

Route of admin.: other: see remarks Exposure period: single administration

Frequency of treatment:

Duration of test: no data

Doses: the highest dose tested was reported to be a saturated acetone

solution corresponding to 0.5 umoles per egg (ca. 173 ug)

Control Group: no data specified

Method:

Year: GLP:

Test substance: other TS: Vulkacit DZ

Remark: the test substance was injected into three day chicken

embryos

Result: Vulkacit DZ did not produce any evidence of embryotoxic

or teratogenic effects

Source: Monsanto

Bayer AG Leverkusen

07-JUN-1994 (24)

5.10 Other Relevant Information

Type: other

Remark: DCBS had no effect on three-day chicken embryos when

injected into the air chamber.

Source: Bayer AG Leverkusen

04-MAY-1992 (25) (26)

5.11 Experience with Human Exposure

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Date: 18-OCT-2001
6. References ID: 4979-32-2

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- (7) de Groot, A.P., CIVO-TNO, short report
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- (17) Monsanto Study ML-88-180
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Date: 18-OCT-2001
6. References ID: 4979-32-2

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- (22) Monsanto Study SR-84-291
- (23) Monsanto Study HL-84-293
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7. Risk Assessment Date: 18-OCT-2001 ID: 4979-32-2

7.1 End Point Summary

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7.2 Hazard Summary

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7.3 Risk Assessment

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IUCLID

Data Set

Existing Chemical ID: 95-31-8 CAS No. 95-31-8

EINECS Name N-tert-butylbenzothiazole-2-sulphenamide

EINECS No. 202-409-1 Molecular Formula C11H14N2S2

Producer Related Part

Company: EUROPEAN COMMISSION - European Chemicals Bureau

Creation date: 11-FEB-2000

Substance Related Part

Company: EUROPEAN COMMISSION - European Chemicals Bureau

Creation date: 11-FEB-2000

Memo: Data for RAPA Sulfenamide Accelerators category

Printing date: 12-OCT-2001 Revision date: 11-FEB-2000 Date of last Update: 11-FEB-2000

Number of Pages: 39

Chapter (profile): Chapter: 1, 2, 3, 4, 5, 7

Reliability (profile): Reliability: without reliability, 1, 2, 3, 4

Flags (profile): Flags: without flag, confidential, non confidential, WGK

(DE), TA-Luft (DE), Material Safety Dataset, Risk

Assessment, Directive 67/548/EEC, SIDS

1. General information 1D. 95-31-8

1.0.1 OECD and Company Information

Name: Akzo Nobel Chemicals b.v. Street: Stationsplein 4, PO Box 247

Town: 3800AE Amersfoort

Country: Netherlands
Phone: +31-33-676767
Telefax: +31-33-676150

Telex: 79322

Name: Bayer Antwerpen N.V.

Street: Haven 507, Scheldelaan 420

Town: Antwerpen Country: Belgium

Name: BFGoodrich Chemical (Belgie) N.V.

Street: Rue de Verdun/straat 742

Town: 1130 Brussels

Country: Belgium

Phone: 32-2-247-19-11 Telefax: 32-2-247-19-91

Name: GENERAL QUIMICA, S.A.

Street: Km.4 Ctra. de Miranda a Puentelarrá Town: 01213 LANTARON COMUNION (ALAVA)

Country: Spain

Phone: 947-31 01 45 Telefax: 947-31 38 88

Telex: 39531

Name: Monsanto Europe N.V. Street: Tervurenlaan, 270-272

Town: 1150 Bruxelles

Country: Belgium

Name: UniroyalChemical Company

Street: Benson Road

Town: 06749 Middlebury, CT

Country: United States
Phone: 203-573-3390
Telefax: 203-573-4531
Telex: 3106710383

1.0.2 Location of Production Site

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1.0.3 Identity of Recipients

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1.1 General Substance Information

Substance type: organic Physical status: solid

1.1.0 Details on Template

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1.1.1 Spectra

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1.2 Synonyms

2-Benzothiazolesulfenamide, N-(1,1-dimethylethyl)-

Source: Akzo Nobel Chemicals b.v. Amersfoort

Monsanto Europe N.V. Bruxelles

2-benzothiazolesulfenamide, N-(1,1-dimethylethyl)-

Remark: other: tertiary-butylbenzothiazolesulfenamide Source: BFGoodrich Chemical (Belgie) N.V. Brussels

Benzothiazolyl-2-tert-butylsulfenamide

Source: GENERAL QUIMICA, S.A. LANTARON COMUNION (ALAVA)

Benzothiazyl-2-tert-butylsulfenamid

Source: Bayer Antwerpen N.V. Antwerpen

Cure-rite BBTS

Source: UniroyalChemical Company Middlebury, CT

Delac NS

Source: UniroyalChemical Company Middlebury, CT

N-(1,1-dimethylethyl)-2-benzothiazolesulfenamide

Source: GENERAL QUIMICA, S.A. LANTARON COMUNION (ALAVA)

N-tert.-butyl-2-benzothiazolsulfenamide

Source: Bayer Antwerpen N.V. Antwerpen

N-tert.butyl-2-benzothiazyl sulphenamide

Source: Akzo Nobel Chemicals b.v. Amersfoort

Perkacit NS

Source: UniroyalChemical Company Middlebury, CT

Santocure NS

Source: UniroyalChemical Company Middlebury, CT

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TBBS

Source: UniroyalChemical Company Middlebury, CT

Akzo Nobel Chemicals b.v. Amersfoort

Bayer Antwerpen N.V. Antwerpen Monsanto Europe N.V. Bruxelles

TBBS/EGC

Source: GENERAL QUIMICA, S.A. LANTARON COMUNION (ALAVA)

tertiary-Butylbenzothiazolesulfenamide

Source: Akzo Nobel Chemicals b.v. Amersfoort

Monsanto Europe N.V. Bruxelles

Vulkacit NZ

Source: UniroyalChemical Company Middlebury, CT

Bayer Antwerpen N.V. Antwerpen

1.3 Impurities

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1.4 Additives

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1.5 Quantity

Quantity 10 000 - 50 000 tonnes

1.6.1 Labelling

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1.6.2 Classification

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1.7 Use Pattern

Type: type

Category: Use in closed system

Type: type

Category: Use resulting in inclusion into or onto matrix

Type: industrial

Category: Chemical industry: used in synthesis

Type: industrial

Category: Polymers industry

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Type: industrial

Category: other: Rubber processing industry

Type: use

Category: Vulcanizing agents

Type: use

Category: other: vulcanisation of natural and sythetic rubber

1.7.1 Technology Production/Use

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1.8 Occupational Exposure Limit Values

Type of limit: TLV (US)
Limit value: 10 mg/m3

Source: GENERAL QUIMICA, S.A. LANTARON COMUNION (ALAVA)

Type of limit: Limit value:

Remark: Occupational exposure limit has not been set.

Source: Akzo Nobel Chemicals b.v. Amersfoort

1.9 Source of Exposure

Source: UniroyalChemical Company Middlebury, CT

Remark: Method of manufacturing: oxidative coupling of

2-mercaptobenzothiazole and t-butylamine using either sodium

hypochlorite or chlorine gas and air.

Source: GENERAL QUIMICA, S.A. LANTARON COMUNION (ALAVA)

1.10.1 Recommendations/Precautionary Measures

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1.10.2 Emergency Measures

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1.11 Packaging

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1.12 Possib. of Rendering Subst. Harmless

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1.13 Statements Concerning Waste

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1.14.1 Water Pollution

Classified by: Labelled by:

Class of danger: 0 (generally not water polluting)

Source: UniroyalChemical Company Middlebury, CT

1.14.2 Major Accident Hazards

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1.14.3 Air Pollution

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1.15 Additional Remarks

Remark: Disposal: Controlled combustion.

Transport information:

UN number: NP ADR/RID: NP IATA-DGR: NP IMDG: NP

Source: GENERAL QUIMICA, S.A. LANTARON COMUNION (ALAVA)

1.16 Last Literature Search

-

1.17 Reviews

_

1.18 Listings e.g. Chemical Inventories

_

- 5/39 -

2. Physico-chemical Data

2.1 Melting Point

Value: ca. 103 degree C

Decomposition: no Sublimation: no Method: other GLP: no data

Source: Monsanto Europe N.V. Bruxelles

(1)

Value: = 104 degree C Method: other: no data

GLP: no data

GENERAL QUIMICA, S.A. LANTARON COMUNION (ALAVA) Source:

Value: ca. 109 degree C

Decomposition: Sublimation: no no Method: other GLP: no data

Source: Monsanto Europe N.V. Bruxelles

(2)

2.2 Boiling Point

Value:

Source: Monsanto Europe N.V. Bruxelles

2.3 Density

density Type:

ca. 1.28 g/cm3 at 25 degree C Value:

Method: other no data GLP:

Source: Monsanto Europe N.V. Bruxelles

(2)

Type: density

Value: = 1.29 g/cm3 at 25 degree C

other: no data Method:

GLP: no data

GENERAL QUIMICA, S.A. LANTARON COMUNION (ALAVA) Source:

density Type: = 1.29 g/cm3Value:

other Method: GLP: no data

Monsanto Europe N.V. Bruxelles Source:

(3)

2.3.1 Granulometry

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Date: 12-OCT-2001 ID: 95-31-8 2. Physico-chemical Data

2.4 Vapour Pressure

Value: .000000611 hPa at 20 degree C

Method: other (measured)

GLP: no data

Source: Monsanto Europe N.V. Bruxelles

(1)

Value: .00000137 hPa at 25 degree C

Method: other (measured)

GLP: no data

Monsanto Europe N.V. Bruxelles Source:

(1)

.0000547 hPa at 25 degree C Value:

Method: other (calculated)

Monsanto Europe N.V. Bruxelles Source:

(1)

2.5 Partition Coefficient

4.38 log Pow:

other (measured) Method:

Year:

GLP: no data

Source: Monsanto Europe N.V. Bruxelles

(4)

log Pow: 4.67 at 22.4 degree C

Method:

1991 Year: GLP: yes

Source: Monsanto Europe N.V. Bruxelles

(1)

2.6.1 Water Solubility

< 1 mg/l at 20 degree C Value:

Method: other GLP: no data

Source: Monsanto Europe N.V. Bruxelles

(1)

Value: .3 mg/lMethod: other GLP: no data

Monsanto Europe N.V. Bruxelles Source:

(5)

Remark: No soluble

GENERAL QUIMICA, S.A. LANTARON COMUNION (ALAVA) Source:

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Date: 12-OCT-2001
2. Physico-chemical Data

ID: 95-31-8

2.6.2 Surface Tension

-

2.7 Flash Point

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2.8 Auto Flammability

_

2.9 Flammability

-

2.10 Explosive Properties

_

2.11 Oxidizing Properties

_

2.12 Additional Remarks

_

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3. Environmental Fate and Pathways

3.1.1 Photodegradation

3.1.2 Stability in Water

abiotic Type:

t1/2 pH7: = 6.1 hour(s)

= 100 % after Degradation: 24 hour(s)

at pH 7

Method: other: ABC Laboratories method

Year: GLP: yes

Test substance: other TS: Monsanto product 1983.

Result: Report indicates that Santocure NS hydrolyzed completely.

Mercaptobenzothiazole and t-butylamine were identified as

hydrolysis products.

Half-life should be viewed as an estimate.

Monsanto Europe N.V. Bruxelles Source:

Test condition: Deionized water filtered through a 0.45 micron filter and

> adjusted to pH 7.00 +/- 0.05 using 0.1 M NaOH/0.1 M KH2PO4 buffer system. Stock solutions were prepared in acetone and

then aliquots transferred to the water system.

Santocure NS supplied by Monsanto Polymer Products Company Test substance:

November 30, 1983. Sample was from lot number NC06-107 with

a purity of 97%.

(6)

Type: abiotic

t1/2 pH7: = 10.4 - 13.2 hour(s)

t1/2 pH9: = 40.5 hour(s)t1/2 pH 5: = 5.1 hour(s)

other: ABC Laboratories Method (1984). Method: 1984 Year: GLP: yes

Test substance: other TS: Monsanto product 1983.

Result: Test results confirm the results obtained in Monsanto report

> AB-84-X128. Santocure NS has a hydrolysis half-life less than 24 hours in water. Presence of sunlight did not impact rate of disappearance of Santocure NS. Data suggest that Santocure NS hydrolyzes to yield mercaptobenzothiazole.

Monsanto Europe N.V. Bruxelles Source:

Deionized and environmental water (well water) were filtered Test condition:

through a 0.45 micron filter. The pH was adjusted to one of 3 pHs. At pH 5 a potassium hydrogen phthalate buffer was used, at pH 7 a potassium dihyrogenphosphate buffer was used while at pH 9 a sodium borate buffer was used. Hydrolysis in sunlight and in the dark and for samples prepared in buffered well water and in buffered deionized water were used to evaluate the impact of sunlight and other minerals on the hydrolysis rate. Neither sunlight nor added minerals

from the well water impacted hydrolysis significantly.

Santocure NS supplied by Monsanto Polymer Products Company Test substance:

November 30, 1983. Sample was from lot number NC06-107

with a purity of 97%.

(7)

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Date: 12-OCT-2001 3. Environmental Fate and Pathways ID: 95-31-8

3.1.3 Stability in Soil

3.2 Monitoring Data (Environment)

3.3.1 Transport between Environmental Compartments

3.3.2 Distribution

3.4 Mode of Degradation in Actual Use

3.5 Biodegradation

Type: aerobic

Type: aeropic
Inoculum: activated sludge
Concentration: 100 mg/l
Degradation: 0 % after 28 day
OFCD Guide-line 3

OECD Guide-line 301 C "Ready Biodegradability: Modified MITI Method:

Test (I)"

Year: 1991 GLP: yes

Test substance: other TS: 99%

Monsanto Europe N.V. Bruxelles Source:

(8)

Type: aerobic
Inoculum: activated sludge, adapted
Concentration: 29.4 mg/l related to Test substance

Result: other: at 32 days 63.5% of theoretical amount of CO2 had

evolved.

other: Monsanto shake flask procedure. Method: 1975 Year: GLP: no data

Test substance: other TS

Source: Monsanto Europe N.V. Bruxelles

Test condition: Monsanto shake flask procedure: 60 mL of acclimated

bacterial seed is mixed with 440 mL of minimal salts media

in fluted 2-liter flask.

Appl. Microbiol. 30:922 (1975).

Monsanto Industrial Chemicals MIC 270582. Test substance:

(9)

3.6 BOD5, COD or BOD5/COD Ratio

- 10/39 -

Date: 12-OCT-2001
3. Environmental Fate and Pathways ID: 95-31-8

3.7 Bioaccumulation

-

3.8 Additional Remarks

-

- 11/39 -

Date: 12-OCT-2001
4. Ecotoxicity ID: 95-31-8

AQUATIC ORGANISMS

4.1 Acute/Prolonged Toxicity to Fish

Type: flow through

Species: Pimephales promelas (Fish, fresh water)

Exposure period: 14 day

Unit: mg/l Analytical monitoring: yes

LC50: > .3

Method: other: Monsanto protocol

Year: 1979 GLP: no data

Test substance: other TS: received from John Vander Kooi

Remark: TOXICTY ABOVE WATER SOLUBILITY (.3 mg/l); 14day LC50>2.33

mg/1

Source: Monsanto Europe N.V. Bruxelles
Test condition: continuous flow diluter; rate=2 ml/1;

solvent=dimethylformamide (.33 mg/l); mean weight=.43 gm;

mean length=38 mm

(10)

Type: static

Species: Brachydanio rerio (Fish, fresh water)

Exposure period: 96 hour(s)

Unit: mg/l Analytical monitoring: yes

Method: other: Directive 67/548/EEC "Acute toxicity to fish" Draft

1992

Year: 1993 GLP: yes

Test substance: other TS: 99%

Remark: No mortality beneath the detection limit of the analytical

method (0.5 mg/ml)

Source: Monsanto Europe N.V. Bruxelles

(8)

Type: static

Species: Lepomis macrochirus (Fish, fresh water)

Exposure period: 96 hour(s)

Unit: mg/l Analytical monitoring: no

LC50: > .3

Method: OECD Guide-line 203 "Fish, Acute Toxicity Test"
Year: 1984 GLP: no data

Test substance: as prescribed by 1.1 - 1.4

Remark: TOXICITY ABOVE WATER SOLUBILITY (.3 mg/l); 96hr LC50=1.2

mg/l (C.I.=.98-1.6 mg/l); 24hr LC50=3.6 mg/l; 48hr

LC50=1.5 mg/1

Source: Monsanto Europe N.V. Bruxelles

Test condition: solvent=acetone; mean length=3.8 cm; 22C

(11)

- 12/39 -

Date: 12-OCT-2001 ID: 95-31-8 4. Ecotoxicity

static

Species: Oncorhynchus mykiss (Fish, fresh water)

Exposure period: 96 hour(s)

Unit: mq/1Analytical monitoring: no

LC50: > .3

Method: OECD Guide-line 203 "Fish, Acute Toxicity Test" 1984 Year: GLP: no data

Test substance: as prescribed by 1.1 - 1.4

TOXICITY ABOVE WATER SOLUBILITY (.3 mg/l); 96hr LC50=1.6 Remark:

mg/l (C.I.=1.3-1.9 mg/l); 24 & 48 hr LC50=1.6 mg/l

Source: Monsanto Europe N.V. Bruxelles
Test condition: solvent=acetone; mean length=3.7 cm; 12C

(11)

Type: static

Species: Pimephales promelas (Fish, fresh water)

Exposure period: 96 hour(s)

Unit: mg/1Analytical monitoring: no

NOEC: > .3 LC50: > .3

OECD Guide-line 203 "Fish, Acute Toxicity Test" Method: Year: 1984 GLP: yes

Test substance: as prescribed by 1.1 - 1.4

TOXICITY ABOVE WATER SOLUBILITY (.3 mg/l); 96 hr LC50=21 Remark:

mg/l (C.I.=6-27 mg/l); 96h NOEC=5.6 mg/l; 24hr LC50=24 mg/l;

48hr LC50=21 mg/l

Source: Monsanto Europe N.V. Bruxelles

Test condition: solvent=acetone; 22C

(12)

4.2 Acute Toxicity to Aquatic Invertebrates

Type:

Species: Daphnia magna (Crustacea)

Exposure period: 48 hour(s)

Unit: mq/1Analytical monitoring: no

NOEC: > .3 EC50: > .3

OECD Guide-line 202, part 1 "Daphnia sp., Acute Method:

Immobilisation Test"

1984 GLP: no data Year:

Test substance: as prescribed by 1.1 - 1.4

Remark: TOXICITY ABOVE WATER SOLUBILITY (0.3 mg/l); 24hr & 48hr

EC50>100 mg/l; 48 hr NOEC=100 mg/l

Monsanto Europe N.V. Bruxelles Source:

Test condition: solvent=acetone

(13)

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4.3 Toxicity to Aquatic Plants e.g. Algae

Species: Selenastrum capricornutum (Algae)

Endpoint: biomass
Exposure period: 96 hour(s)

Unit: mg/l Analytical monitoring: no

EC50: > .3

Method: other: U.S. EPA (1971) Algal Assay Procedure: Bottel Test

EPA1972-795-146/1

Year: 1971 GLP: no data

Test substance: as prescribed by 1.1 - 1.4

Remark: TOXICITY ABOVE WATER SOLUBILITY (.3 mg/l); 96hr EC50=4 mg/l

(C.I.=1-15 mg/l); in vivo chlorophyll results: 24 hr EC50>10 $\,$

mg/l, 48hr EC50=6 mg/l, 72 & 96hr EC50=4 mg/l

Source: Monsanto Europe N.V. Bruxelles

Test condition: solvent=acetone; 24C; 4000 lux, 10,000 cells/ml

(14)

4.4 Toxicity to Microorganisms e.g. Bacteria

Type: aquatic

Species: activated sludge

Exposure period: 3 hour(s)

Unit: mg/l Analytical monitoring: no

EC50: > 10000

Method: ISO 8192 "Test for inhibition of oxygen consumption by

activated sludge"

Year: 1990 GLP: yes

Test substance: other TS: 99%

Source: Monsanto Europe N.V. Bruxelles

(8)

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- 4.5 Chronic Toxicity to Aquatic Organisms
- 4.5.1 Chronic Toxicity to Fish

-

4.5.2 Chronic Toxicity to Aquatic Invertebrates

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TERRESTRIAL ORGANISMS

4.6.1 Toxicity to Soil Dwelling Organisms

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4.6.2 Toxicity to Terrestrial Plants

-

4.6.3 Toxicity to other Non-Mamm. Terrestrial Species

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4.7 Biological Effects Monitoring

_

4.8 Biotransformation and Kinetics

_

4.9 Additional Remarks

_

- 15/39 -

Date: 12-OCT-2001 ID: 95-31-8 5. Toxicity

5.1 Acute Toxicity

5.1.1 Acute Oral Toxicity

LD50 Type: Species: rat

Strain: Sex: Number of Animals: Vehicle:

Value: > 6310 mg/kg bw

other: Younger Laboratory method Method:

Year: 1973 GLP: no data

Test substance: as prescribed by 1.1 - 1.4

TEST CONDITIONS: Remark:

> mode of administration: gavage number of animals: 5/dose (M,F)

TEST RESULTS: lung congestion; liver, discoloration; GIT, inflammation; apetite, decr; activity, decr; weakness;

collapse.

Source: Monsanto Europe N.V. Bruxelles

(15)

Type: LD50 Species: rat

Strain: Sex: Number of Animals: Vehicle:

Value: = 6850 mg/kg bw

Method: other

GLP: no data Year:

Test substance: as prescribed by 1.1 - 1.4

Remark: TEST CONDITIONS:

number of animals: 2 M, 3 F

vehicle: water (10% w/w suspension)

TEST QUALITY:

too few animals used

TEST RESULTS: mortality

Source: Monsanto Europe N.V. Bruxelles

(16)

- 16/39 -

Type: LD0 Species: rat

Strain: Sex: Number of Animals: Vehicle:

> 10000 mg/kg bw Value:

Method: other

Year: GLP: no data

Test substance: as prescribed by 1.1 - 1.4

Remark: TEST CONDITIONS:

number of animals: 1 M, screening test

doses: one dose level

vehicle: water (25 % w/w suspension)

TEST RESULTS: no mortality

Source: Monsanto Europe N.V. Bruxelles

(17)

Type: LDLo Species: rat

Strain: Sex: Number of Animals: Vehicle:

Value: = 7940 mg/kg bw Method: other: no data

Year: 1990 GLP: no data

Test substance: as prescribed by 1.1 - 1.4 Source: GENERAL OUIMICA, S.A. LANT GENERAL QUIMICA, S.A. LANTARON COMUNION (ALAVA)

(18)

5.1.2 Acute Inhalation Toxicity

5.1.3 Acute Dermal Toxicity

LD50 Type: Species: rabbit

Strain: Sex: Number of Animals: Vehicle:

Value: > 7940 mg/kg bw

Method: other: Younger Laboratories method

Year: 1973 GLP: no data

Test substance: as prescribed by 1.1 - 1.4

Remark: TEST CONDITIONS:

mode of administration: not given

number of animals: 1 M, 1 F

- 17/39 -

exposure time: 24 hours

vehicle: 40% suspension in corn oil

Source: Monsanto Europe N.V. Bruxelles

(15)

Type: LD50 Species: rabbit

Strain:
Sex:
Number of
Animals:
Vehicle:

Value: < 7940 mg/kg bw Method: other: no data

Year: 1990 GLP: no data

Test substance: as prescribed by 1.1 - 1.4

Source: GENERAL QUIMICA, S.A. LANTARON COMUNION (ALAVA)

(18)

Type: LD0 Species: rabbit

Strain:
Sex:
Number of
Animals:
Vehicle:

Value: > 6000 mg/kg bw

Method: other

Year: GLP: no data

Test substance: as prescribed by 1.1 - 1.4

Remark: TEST CONDITIONS:

number of animals: 1 M, 1 F/dose

TEST RESULTS:

all animals survived at 6000 mg/kg bw

Source: Monsanto Europe N.V. Bruxelles

(19)

5.1.4 Acute Toxicity, other Routes

Type: LD50 Species: mouse

Strain:
Sex:
Number of
 Animals:
Vehicle:

Route of admin.: i.p.

Value: = 5000 mg/kg bw Method: other: no data

Year: 1976 GLP: no data

Test substance: as prescribed by 1.1 - 1.4

Source: GENERAL QUIMICA, S.A. LANTARON COMUNION (ALAVA)

(20)

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Date: 12-OCT-2001 ID: 95-31-8 5. Toxicity

LC50 Type: Species: mouse

Strain: Sex: Number of Animals: Vehicle:

Route of admin.: i.p.

= 5000 mg/kg bwValue:

Method:

GLP: no data

= 5000 other
Year: Year: 1976 Test substance: as prescribed by 1.1 - 1.4 Source: Monsanto Europe N.V. Bruxelles

(21)

LD50 Type: Species: mouse

Strain: Sex: Number of Animals: Vehicle:

Route of admin.: i.v.

= 180 mg/kg bw other: no data Value: Method:

Year: GLP: no data

Test substance: as prescribed by 1.1 - 1.4

Source: GENERAL QUIMICA, S.A. LANTARON COMUNION (ALAVA)

(22)

Type: LC50 Species: mouse

Strain: Sex: Number of Animals: Vehicle:

Route of admin.: i.v.

= 180 mg/kg bwValue:

Method: other

Year: GLP: no data

Test substance: as prescribed by 1.1 - 1.4 Source: Monsanto Europe N.V. Bruxelles

(23)

- 19/39 -

5.2 Corrosiveness and Irritation

5.2.1 Skin Irritation

Species: rabbit

Concentration:

Exposure:
Exposure Time:
Number of
Animals:
PDII:

Result: not irritating EC classificat.: not irritating

Method: other: Younger Laboratories method

Year: 1973 GLP: no data

Test substance: as prescribed by 1.1 - 1.4

Remark:

TEST CONDITIONS:

mode of administration: as per FHSA

exposure time: 24 hours

reading times: 24, 48, 72, 168 hours

vehicle: water
skin: intact

number of animals: 6

TEST RESULTS:

EU mean erythema score: 0.0 EU mean edema score: 0.0

PII: 0/8

Source: Monsanto Europe N.V. Bruxelles

(15)

Species: rabbit

Concentration:

Exposure:
Exposure Time:
Number of
Animals:

PDII:

Result: slightly irritating EC classificat.: not irritating

Method: other: modified Draize

Year: 1944 GLP: no data

Test substance: as prescribed by 1.1 - 1.4

Remark: TEST CONDITIONS:

number of animals: 1 M, 2 F exposure time: 24 hours

PII: 1.0/8

healing time: 2 hours

Source: Monsanto Europe N.V. Bruxelles

(17)

- 20/39 -

Species: rabbit

Concentration:

Exposure:
Exposure Time:
Number of
Animals:
PDII:

Result: slightly irritating

EC classificat.: not irritating

Method: other: modified Draize

Year: 1944 GLP: no data

Test substance: as prescribed by 1.1 - 1.4

Remark: TEST CONDITIONS:

number of animals: 6 M exposure time: 24 hours

skin: intact

TEST RESULTS: PII: 1.5/8.0

healing time: not cleared in 72 hours

Source: Monsanto Europe N.V. Bruxelles

(24)

5.2.2 Eye Irritation

Species: rabbit

Concentration:

Dose:

Exposure Time:
Comment:
Number of
Animals:

Result: slightly irritating EC classificat.: not irritating Method: Draize Test

Year: 1944 GLP: no data

Test substance: as prescribed by 1.1 - 1.4

Remark: TEST CONDITIONS: number of animals: 6

TEST RESULTS:

EU mean erythema score: 2.53 EU mean chemosis score: 0 EU mean corneal score: 0 EU mean iritis score: 0 Draize score: 2.5/110

Source: Monsanto Europe N.V. Bruxelles

(15)

- 21/39 -

Date: 12-OCT-2001 ID: 95-31-8 5. Toxicity

Species: rabbit

Concentration:

Dose:

Exposure Time: Comment:

Number of Animals:

Result: slightly irritating

EC classificat.: not irritating

Method: Year: other: modified Draize

1944 GLP: no data

Test substance: as prescribed by 1.1 - 1.4

Remark: TEST CONDITIONS:

number of animals: 2 M, 1 F

TEST RESULTS:

Draize score: 2.4/110 healing time: 72 hours

Source: Monsanto Europe N.V. Bruxelles

(17)

Species: rabbit

Concentration:

Dose:

Exposure Time: Comment: Number of Animals:

Result: slightly irritating EC classificat.: not irritating Method: Draize Test

GLP: no data Year: 1944

Test substance: as prescribed by 1.1 - 1.4

TEST CONDITIONS: Remark:

number of animals: 6 M

TEST RESULTS:

Draize score: 2.5/110.0 healing time: 48 hours

Monsanto Europe N.V. Bruxelles Source:

(25)

- 22/39 -

5.3 Sensitization

Type: Buehler Test Species: guinea pig

Number of
Animals:
Vehicle:

Result: sensitizing Classification: sensitizing

Method: other: Pharmakon Laboratory method
Year: 1980 GLP: yes

Test substance: as prescribed by 1.1 - 1.4

Remark: TEST CONDITIONS:

The test material came from three different locations. It

was applied in a 25% preparation in ethanol.

Source: Monsanto Europe N.V. Bruxelles

(26)

Type: Patch-Test Species: human

Number of
Animals:
Vehicle:

Result: sensitizing Classification: sensitizing

Method: other Year: 1969

Year: 1969 GLP: no data

Test substance: as prescribed by 1.1 - 1.4

Remark: 55 subjects (18 M/ 37 F); 24 hour occluded, alternate day

patches for 14 days.

9 Of 55 showed sensitization; not a primary irritant

Source: Monsanto Europe N.V. Bruxelles

(27)

Type: Patch-Test Species: human

Number of
Animals:
Vehicle:

Result: sensitizing Classification: sensitizing

Method: other: Not specified

Year: 1983 GLP: no data

Test substance: as prescribed by 1.1 - 1.4

Remark: TEST CONDITIONS:

1% Santocure NS in petrolatum. No testing was carried out

to determine if trace quantities of MBT were in the

preparation.
TEST RESULTS:

13 of 14 subjects who had previously been sensitized by MBT also showed sensitivity reactions when exposed to Santocure

NS.

Source: Monsanto Europe N.V. Bruxelles

(28)

- 23/39 -

Type: Patch-Test Species: human

Number of Animals: Vehicle:

Result: sensitizing Classification: sensitizing

Method: other: Product Investigations method Year: 1982 GLP: no data

Test substance: as prescribed by 1.1 - 1.4

Remark: TEST CONDITIONS:

60% preparation in petrolatum.

TEST RESULTS:

13 of 45 subjects responded during challenge.

Source: Monsanto Europe N.V. Bruxelles

(29)

Type: other Species: rabbit

Number of
Animals:
Vehicle:

Result: not sensitizing Classification: not sensitizing

Method: other: Pharmakon Laboratory method
Year: 1982 GLP: yes

Test substance: as prescribed by 1.1 - 1.4

Remark: The studies were comedogenicity assays.

Three separate samples of Santocure NS were tested in concentrations of 0.01, 0.1, or 10% in chloroform (sample #1), 1, 10, or 100% in chloroform (sample #2), and 0.01,

0.1, or 1% in chloroform (sample #3).

Source: Monsanto Europe N.V. Bruxelles

(30)

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5.4 Repeated Dose Toxicity

Species: rat Sex: male/female

Strain: Sprague-Dawley Route of admin.: inhalation Exposure period: 4 weeks

Frequency of

treatment: 6 hours/day, 5 days/week

Post. obs.

period: no

Doses: 0.0, 0.0024, 0.029, and 0.084 mg/l

Control Group: yes

NOAEL: = .029 mg/l

Method: other: Monsanto Laboratory method Year: 1978 GLP: yes

Test substance: as prescribed by 1.1 - 1.4

Result: 1) 0.0024 mg/l:

no effects.
2) 0.029 mg/l:

NOAEL;

blood, SGOT, incr.
3) 0.084 mg/l:
blood, SGOT, incr;
body weight, decr;
liver, histopath;

lymph nodes, histopath.

Source: Monsanto Europe N.V. Bruxelles

(31)

Species: rat Sex: male/female

Strain: Sprague-Dawley
Route of admin.: oral feed
Exposure period: 4 weeks

Frequency of

treatment: daily

Post. obs.

period: none

Doses: 0, 10, 50, 300, 1000, 3000 mg/kg bw/day

Control Group: yes

NOAEL: 1000 mg/kg bw LOAEL: 3000 mg/kg bw

Method: other

Year: 1978 GLP: yes

Test substance: as prescribed by 1.1 - 1.4

Remark: TEST CONDITIONS:

number of animals: 5 M, 5 F/dose

Result: 1) 1000 mg/kg bw/d:

no effect;

2) 3000 mg/kg bw/d: body weight, decr; food consumption, decr;

LOAEL.

Source: Monsanto Europe N.V. Bruxelles

(32)

Species: rat Sex: male/female

Strain: Sprague-Dawley

Route of admin.: gavage Exposure period: 30 days

Frequency of

treatment: daily

Post. obs.

period: none

Doses: 0, 100, 300, 1000, and 3000 mg/kg bw/d

Control Group: yes

LOAEL: = 100 mg/kg bw

Method: other: Bio/dynamics Laboratory method Year: 1981 GLP: yes

Test substance: as prescribed by 1.1 - 1.4

Remark: All female rats in the 3000 mg/kg bw/d group and 1 male rat

in the 3000 mg/kg bw/d group died by day 6. All other 3000

mg/kg bw/day male rats were sacrificed on day 6 in a

moribund condition. TEST CONDITIONS:

number of animals: 5/sex/group

Result: 1) 100 mg/kg bw/d:

LOAEL;

heart weight, decr, F.

2) 300 mg/kg bw/d:
body weight, decr, M;
heart, weight, decr, F;
kidney, weight, incr;
liver, weight, incr, F.

3) 1000 mg/kg bw/d:
body, weight, decr, M;
heart, weight, decr;
kidney, weight, incr;
liver, weight, incr, F.

4) 3000 mg/kg bw/d:

mortality.

Source: Monsanto Europe N.V. Bruxelles

(33)

- 26/39 -

Species: rat Sex: male/female

Strain: Sprague-Dawley

Route of admin.: gavage Exposure period: 90 d

Frequency of

treatment: daily

Post. obs.

period: no

Doses: 0, 100, 300, and 1000 mg/kg bw/d

Control Group: yes

NOAEL: = 100 mg/kg bwLOAEL: = 300 mg/kg bw

Method: other: Monsanto Laboratory method
Year: 1982 GLP: yes

Test substance: as prescribed by 1.1 - 1.4

Remark: TEST CONDITIONS:

number of animals: 5/sex/group

Result: 1) 100 mg/kg bw/d:

NOEL, NOAEL; no effects.

2) 300 mg/kg bw/d:

LOAEL;

body, weight, decr, M.
3) 1000 mg/kg bw/d:
body, weight, decr, M;
liver, weight, incr, F;
kidney, weight, incr, F;
serum, cholesterol, incr, F;
urine, specific gravity, incr, F.

Source: Monsanto Europe N.V. Bruxelles

(34)

Species: rat Sex: male

Strain: Sprague-Dawley

Route of admin.: gavage Exposure period: 1-3 days

Frequency of

treatment: multiple daily doses (varied)

Post. obs.

period: up to 14 days

Doses: up to 40 grams total dose

Control Group: no

NOAEL: > 40000 mg/kg bw

Method: other: screen 1 male/ dose

Year: GLP: no data

Test substance: as prescribed by 1.1 - 1.4

Remark: TEST CONDITIONS:

number of animals: 1 M/dose

doses: ranged from 15 to 40 grams/kg bw.

Source: Monsanto Europe N.V. Bruxelles

(17)

- 27/39 -

Species: rat Sex: male/female

Strain: Sprague-Dawley

Route of admin.: gavage Exposure period: 1-5 days

Frequency of

treatment: multiple daily doses

Post. obs.

period: up to 14 days

Doses: up to 6000 mg/kg bw/d

Control Group: no

NOAEL: > 6000 mg/kg bw

Method: other

Year: GLP: no data

Test substance: as prescribed by 1.1 - 1.4

Remark: TEST CONDITIONS:

1-2 males or females dosed with multiple daily doses for up to 5 days and up to a total of 60 grams/kg of test material.

TEST RESULTS:

all animals survived

Source: Monsanto Europe N.V. Bruxelles

(25)

Species: rabbit Sex: male/female

Strain: New Zealand white

Route of admin.: dermal Exposure period: 21 d

Frequency of

treatment: daily

Post. obs.

period: no

Doses: 0, 125, 500, or 2000 mg/kg bw/d

Control Group: yes

NOAEL: > 2000 mg/kg bw

Method: other: IRDC Laboratory method

Year: 1979 GLP: yes

Test substance: as prescribed by 1.1 - 1.4

Remark: TEST CONDITIONS:

number of animals: 10/sex/group

Result: 1) 125 mg/kg bw/d:

skin, irritation, slight.
2) 500 mg/kg bw/d:
skin, irritation, slight.

3) 2000 mg/kg bw/d:

NOAEL;

skin, irritation, slight.

Source: Monsanto Europe N.V. Bruxelles

(35)

- 28/39 -

5.5 Genetic Toxicity 'in Vitro'

Type: Ames test

System of

testing: Salmonella typhimurium strains TA98, 100, 1535, 1537, 1538

Concentration:
Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative Method: other

Year: 1975 GLP: no data

Test substance: as prescribed by 1.1 - 1.4
Source: Monsanto Europe N.V. Bruxelles

(36)

Type: Ames test

System of

testing: Salmonella typhimurium strains TA98, 100, 1535, 1537, 1538

Concentration:
Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative Method: other

Year: 1982 GLP: no data

Test substance: as prescribed by 1.1 - 1.4
Source: Monsanto Europe N.V. Bruxelles

(37)

Type: Ames test

System of

testing: Salmonella typhimurium strains TA98, 100, 1535, 1537, 1538

Concentration: up to 500 ug/plate

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method: other: Bionetics method

Year: 1975 GLP: no data

Test substance: as prescribed by 1.1 - 1.4
Source: Monsanto Europe N.V. Bruxelles

(38)

- 29/39 -

Type: Ames test

System of

testing: Salmonella typhimurium strains TA98, 100, 1535, 1537, 1538

Concentration: up to 5000 ug/plate

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method: other: Bionetics method

Year: 1975 GLP: yes

Test substance: as prescribed by 1.1 - 1.4
Source: Monsanto Europe N.V. Bruxelles

(39)

Type: Bacterial gene mutation assay

System of

testing: Salmonella typhimurium TA 98, 100, 1535, 1537, 1538

Concentration: up to 3000 ug/plate

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method: other: no data

Year: GLP: no data

Test substance: no data

Source: Monsanto Europe N.V. Bruxelles

(40)

Type: DNA damage and repair assay

System of

testing: E. coli WP2uvrA-(WU-)
Concentration: up to 1000 ug/plate

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: ambiguous

Method: other: Bionetics method

Year: 1956 GLP: yes

Test substance: as prescribed by 1.1 - 1.4 Source: Monsanto Europe N.V. Bruxelles

(39)

Type: Escherichia coli reverse mutation assay

System of

testing: E. coli W3110(polA+), p3078(polA-)

Concentration: up to 1000 ug/plate

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method: other: Bionetics method

Year: 1971 GLP: yes

Test substance: as prescribed by 1.1 - 1.4 Source: Monsanto Europe N.V. Bruxelles

Source: Monsanto Europe N.V. Bruxelles (39)

- 30/39 -

Type: HGPRT assay

System of

testing: CHO cells

Concentration:
Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative Method: other

Year: 1984 GLP: yes

Test substance: as prescribed by 1.1 - 1.4
Source: Monsanto Europe N.V. Bruxelles

(41)

Type: Mammalian cell gene mutation assay

System of

testing: CHO cells
Concentration: up to 10 ug/ml

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method: other: Bionetics method

Year: 1961 GLP: yes

Test substance: as prescribed by 1.1 - 1.4
Source: Monsanto Europe N.V. Bruxelles

(39)

Type: Mammalian cell gene mutation assay

System of

testing: BALB/3T3 cells Concentration: up to 35 ug/ml

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method: other: Bionetics method

Year: 1987 GLP: yes

Test substance: as prescribed by 1.1 - 1.4

Remark: TEST CONDITIONS:

cytotoxicity was observed at a 46% rate at 35 ug/ml.

Source: Monsanto Europe N.V. Bruxelles

(39)

- 31/39 -

Type: Mammalian cell gene mutation assay

System of testing:

Concentration:

40 mg/l

Cytotoxic Conc.:

Metabolic

activation: with

Result:

Method: other: no data

Year: 1983 GLP: no data

Test substance: as prescribed by 1.1 - 1.4

Source: GENERAL QUIMICA, S.A. LANTARON COMUNION (ALAVA)

(42)

Type: Mammalian cell gene mutation assay

System of testing:

Concentration: 35 mg/l

Cytotoxic Conc.:

Metabolic

activation: with

Result:

Method: other: no data

Year: 1983 GLP: no data

Test substance: as prescribed by 1.1 - 1.4

Source: GENERAL QUIMICA, S.A. LANTARON COMUNION (ALAVA)

(42)

Type: Mouse lymphoma assay

System of

testing: L5178Y

Concentration: up to 15 ug/ml without S9 and 60 ug/ml with S9

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: positive

Method: other: Bionetics method

Year: 1975 GLP: yes

Test substance: as prescribed by 1.1 - 1.4

Remark: Negative without S9.

Source: Monsanto Europe N.V. Bruxelles

(39)

- 32/39 -

Type: Mouse lymphoma assay

System of

testing: L5178Y

Concentration: up to 12.5 ug/ml without S9 and 50 ug/ml with S9

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: positive

Method: other: Bionetics method

Year: 1975 GLP: yes

Test substance: as prescribed by 1.1 - 1.4

Remark: Negative without S9.

Source: Monsanto Europe N.V. Bruxelles

(43)

Type: Mouse lymphoma assay

System of

testing: L5178Y

Concentration: up to 100 ug/ml

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: positive

Method: other: SRI method

Year: 1978 GLP: yes

Test substance: as prescribed by 1.1 - 1.4

Remark: Negative without S9.

Source: Monsanto Europe N.V. Bruxelles

(44)

Type: Mouse lymphoma assay

System of

testing: L5178Y

Concentration: up to 100 ug/ml

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: positive

Method: other: SRI method

Year: 1978 GLP: yes

Test substance: as prescribed by 1.1 - 1.4 Source: Monsanto Europe N.V. Bruxelles

(45)

- 33/39 -

Type: Yeast gene mutation assay

System of

testing: S. cerevisiae strain D4 Concentration: up to 500 ug/plate

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method: other: Bionetics method

Year: 1976 GLP: no data

Test substance: as prescribed by 1.1 - 1.4
Source: Monsanto Europe N.V. Bruxelles

(38)

5.6 Genetic Toxicity 'in Vivo'

-

5.7 Carcinogenicity

-

5.8 Toxicity to Reproduction

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5.9 Developmental Toxicity/Teratogenicity

Species: rat Sex: female

Strain: Sprague-Dawley

Route of admin.: gavage

Exposure period: days 6-15 of gestation

Frequency of

treatment: daily
Duration of test: no

Doses: 0, 50, 150, and 500 mg/kg bw/d

Control Group: yes

NOAEL Maternalt.: > 500 mg/kg bw
NOAEL Teratogen.: > 500 mg/kg bw
Method: other: IRDC method

Year: 1978 GLP: yes

Test substance: as prescribed by 1.1 - 1.4

Result: 1) 50 mg/kg bw/d:

no effects.

2) 150 mg/kg bw/d:

no effects.

3) 500 mg/kg bw/d:

maternal:
NOEL, NOAEL;
offspring:
NOEL, NOAEL;
no effects.

Source: Monsanto Europe N.V. Bruxelles

(46)

Species: other Sex: female

Strain: Leghorn

Route of admin.:
Exposure period:
Frequency of
 treatment:
Duration of test:

Doses:

Control Group:

Method:

Year: GLP:

Test substance:

Remark: TBBS has been reported to be embryotoxic and teratogenic in

chick embryos; however, the relevance of this system to man

is questionable.

Source: Monsanto Europe N.V. Bruxelles

(47)

Species: other Sex: female

Strain: Leghorn

Route of admin.:
Exposure period:
Frequency of
treatment:
Duration of test:

Doses:

Control Group:

Method:

Year: GLP:

Test substance:

Remark: TBBS has been reported to be embryotoxic and teratogenic in

chick embryos; however, the relevance of this system to man

is questionable.

Source: Monsanto Europe N.V. Bruxelles

(48)

5.10 Other Relevant Information

-

5.11 Experience with Human Exposure

Remark: There are no reports from manufacturing facilities or the

customer base which suggest that the experimental sensitizing potential is a problem in practice.

Source: Monsanto Europe N.V. Bruxelles

- 35/39 -

Date: 12-OCT-2001 6. References ID: 95-31-8

- (1) Bayer AG Data
- (2) Monsanto Product Specification
- (3) Safety Data Sheet Bayer AG
- (4) Monsanto Study MO-92-9052
- (5) Monsanto Study MO 92-9052
- (6) Monsanto report AB-84-X128.
- (7) Monsanto report AB-84-X133.
- (8) Bayer AG data
- (9) Monsanto report ES-78-SS28.
- (10) Monsanto report ES-79-SS22
- (11) Monsanto report BN-76-0173
- (12) Monsanto report AB-79-0321
- (13) Monsanto report AB-78-0119
- (14) Monsanto report BN-78-0364
- (15) Monsanto Study No. Y-73-222.
- (16) Monsanto study SA-44, April 1954, Scientific Associates Lab.
- (17) Monsanto study SA-85, Dec. 1954, Scientific Associates Lab.
- (18) RTECS: "Registry of Toxic Effects of Chemical Substances" Acute Toxicity Data (ATDAEI): 1, 104, 1990
- (19) Monsanto study SA-85, Dec. 1954 Scientific Associates Lab.
- (20) RTECS: "Registry of Toxic Effects of Chemical Substances" International Polymer Science and Technology (IPSTB3): 3, 93, 1976
- (21) International Polymer Science Tochnology 3, 93. 1976. Reported in RTECS.
- (22) RTECS: "Registry of Toxic Effects of Chemical Substances" U.S. Army Armament Research & Development Command, Chemical Systems Laboratory, NIOSH Exchange Chemicals. (Aberdeen Proving Ground, MD 21010) NX#02241 (CSLNX)

- 36/39 -

Date: 12-OCT-2001
6. References ID: 95-31-8

o. Relefences

(23) U. S. Army. No date. U. S. Army Armament Research and Development Command. Chemical Systems Laboratory, NIOSH Exchange Chemicals. NY Nos. 02241, 02243, and 02251. As reported in RTECS.

- (24) Monsanto study SA-44, Apr 1954, Scientific Associates Lab.
- (25) Monsanto study SA-44, Apr. 1954, Scientific Associates Lab.
- (26) Monsanto Study Nos. PK-82-41 & 47.
- (27) Monsanto study SH-69-10; Food and Drug Research Lab.
- (28) Foussereau, J., Menezes-Brandao, F., Cavelier, C., and Herve-Bazin, B. 1983. Allergy to MBT and its derivatives. Contact Dermatitis 9, 514-516.
- (29) Monsanto Study No. SH-82-11.
- (30) Monsanto Study Nos. PK-82-42 & 48.
- (31) Monsanto Study No. IR-78-95.
- (32) Monsanto study IR-78-195
- (33) Monsanto Study No. BD-81-330.
- (34) Monsanto Study No. ML-82-253.
- (35) Monsanto Study No. IR-79-263.
- (36) Hinderer, R. K., Myhr, B., Jagnnath, D. R., Galloway, S. M., Mann, S. W., Riddle, J. C., and Brusick, D. J. 1983. Mutagenic evaluations of four rubber accelerators in a battery of in vitro mutagenic assays. Environ. Mutagen. 5, 193-215.
- (37) You, X., Zhou, Y., and Hu, Y. 1982. Mutagenicity of fourteen rubber accelerators. Huanjing Kexue 3, 39-42 (as reported in Chem. Abstr. 98:84705t).
- (38) Monsanto Study No. BO-76-181.
- (39) Monsanto Study No. XX-87-9007 (B. F. Goodrich report).
- (40) Sumitomo Chemical Co. data, September 24, 1981 (reported to Bayer AG)
- (41) B. F. Goodrich. 1984. Evaluation of TBBS (BBTS) in the CHO/HGPRT assay.

- 37/39 -

Date: 12-OCT-2001
6. References ID: 95-31-8

(42) RTECS: "Registry of Toxic Effects of Chemical Substances" Environmental Mutagenesis (ENMUDM): 5, 193, 1983

- (43) Monsanto Study No. BO-78-222.
- (44) Monsanto Study No. SR-81-46.
- (45) Monsanto Study No. SR-81-47.
- (46) Monsanto Study No. IR-78-101.
- (47) Korhonen, A., Hemminki, K., and Vaino, H. 1982. Embryotoxicity of benzothiazoles, benzenesulfohydrazine and dithiodimorpholine to the chicken embryo. Arch. Environ. Contam. Toxicol. 11, 753-759.
- (48) Korhonen, A., Hemminki, K., and Vaino, H. 1983. Toxicity of rubber chemicals toward three-day chicken embryos. Scand. J. Work Environ. Health 9, 115-119.

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7. Risk Assessment Date: 12-OCT-2001 ID: 95-31-8

7.1 End Point Summary

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7.2 Hazard Summary

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7.3 Risk Assessment

-

- 39/39 -

IUCLID

Data Set

Existing Chemical ID: 95-33-0 CAS No. 95-33-0

EINECS Name N-cyclohexylbenzothiazole-2-sulphenamide

EINECS No. 202-411-2

TSCA Name 2-Benzothiazolesulfenamide, N-cyclohexyl-

Molecular Formula C13H16N2S2

Producer Related Part

Company:

Creation date: 10-JAN-2000

Substance Related Part

Company:

Creation date: 10-JAN-2000

Memo: Data for RAPA Sulfenamide Accelerators category

Printing date: 17-OCT-2001

Revision date:

Date of last Update: 17-OCT-2001

Number of Pages: 45

Chapter (profile): Chapter: 1, 2, 3, 4, 5, 7

Reliability (profile): Reliability: without reliability, 1, 2, 3, 4

Flags (profile): Flags: without flag, confidential, non confidential, WGK

(DE), TA-Luft (DE), Material Safety Dataset, Risk

Assessment, Directive 67/548/EEC, SIDS

Date: 17-OCT-2001

1. General Information ID: 95-33-0

1.0.1 OECD and Company Information

_

1.0.2 Location of Production Site

_

1.0.3 Identity of Recipients

_

1.1 General Substance Information

Substance type: organic Physical status: solid

Purity: >= 96 % w/w

Remark: cooperating companies:

Bayer Antwerpen N.V., Belgium AKZO Chemicals, Netherlands

Manufacture Landaise de Produits Chimiques, France

Monsanto Europe N.V., Belgium

Uniroyal Chemical Limited, United Kingdom

Source: Bayer AG Leverkusen

27-MAY-1994

1.1.0 Details on Template

-

1.1.1 Spectra

_

1.2 Synonyms

CBS

03-JUN-1994

N-CYCLOHEXYL-2-BENZOTHIAZOLESULFENAMIDE

03-APR-1992

Santocure accelerator

15-OCT-2001

1.3 Impurities

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1.4 Additives

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- 1/45 -

Date: 17-OCT-2001

1. General Information ID: 95-33-0

1.5 Quantity

_

1.6.1 Labelling

_

1.6.2 Classification

_

1.7 Use Pattern

Type: type

Category: Use resulting in inclusion into or onto matrix

15-OCT-2001

Type: industrial

Category: Polymers industry Source: Bayer AG Leverkusen

26-APR-1994

Type: use

Category: Vulcanizing agents Source: Bayer AG Leverkusen

26-APR-1994

1.7.1 Technology Production/Use

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1.8 Occupational Exposure Limit Values

_

1.9 Source of Exposure

_

1.10.1 Recommendations/Precautionary Measures

_

1.10.2 Emergency Measures

_

1.11 Packaging

_

1.12 Possib. of Rendering Subst. Harmless

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- 2/45 -

Date: 17-OCT-2001

1. General Information

ID: 95-33-0

1.13 Statements Concerning Waste

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1.14.1 Water Pollution

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1.14.2 Major Accident Hazards

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1.14.3 Air Pollution

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1.15 Additional Remarks

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1.16 Last Literature Search

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1.17 Reviews

-

1.18 Listings e.g. Chemical Inventories

- 3/45 -

2. Physico-chemical Data

2.1 Melting Point

Value: 93 - 100 degree C

Decomposition: Sublimation: no

Method: other: Handbook value

GLP: no

Testsubstance: other TS: purity not noted Reliability: (2) valid with restrictions

Data from Handbook or collect

Data from Handbook or collection of data

Critical study for SIDS endpoint Flaq:

15-OCT-2001 (1)

< 96 degree C Value:

Source: Bayer AG Leverkusen

22-SEP-2000 (2)

2.2 Boiling Point

Value:

Decomposition: yes

Remark: n.a., product decomposes during destillation

Source: Bayer AG Leverkusen

18-APR-2001

2.3 Density

density Type:

Value: 1.27 g/cm3 at 25 degree C other: Handbook value Method: Testsubstance: other TS: purity not noted (2) valid with restrictions Reliability:

Data from Handbook or collection of data

Critical study for SIDS endpoint Flaq:

15-OCT-2001 (1)

relative density Type:

1.3 g/cm3 at 25 degree C Value: Source: Bayer AG Leverkusen

22-SEP-2000 (2)

2.3.1 Granulometry

2.4 Vapour Pressure

< .0000004 hPa at 25 degree C Value:

Method: other (measured)

Reliability: (2) valid with restrictions

22-SEP-2000 (3)

- 4/45 -

2. Physico-chemical Data

.000000015 hPa at 20 degree C Value:

Source: Bayer AG Leverkusen

16-SEP-1993 (2)

Value: .000000038 hPa at 25 degree C

Source: Bayer AG Leverkusen

16-SEP-1993 (2)

2.5 Partition Coefficient

log Pow:

Method: other (calculated)

Year:

(2) valid with restrictions Reliability:

Accepted calculation method

Critical study for SIDS endpoint Flag:

(4)15-OCT-2001

log Pow: 4.93

Method: other (measured)

Year:

Reliability: (2) valid with restrictions

15-OCT-2001 (3)

2.6.1 Water Solubility

.24 other: ppm at 21 degree C

Qualitative: not soluble

:Hq

Source: Bayer AG Leverkusen

18-APR-2001 (5) (6)

Value: .32 other: ppm at 21 degree C

Qualitative: not soluble

pH: 7

Bayer AG Leverkusen Source:

(5) (6) 18-APR-2001

Value: .48 other: ppm at 21 degree C

Qualitative: .48 other: g

:Hq 9

Source: Bayer AG Leverkusen

Reliability: (2) valid with restrictions

(5) (6) 18-APR-2001

2.6.2 Surface Tension

- 5/45 -

Date: 17-OCT-2001
2. Physico-chemical Data

ID: 95-33-0

2. Figs160 chemical baca

2.7 Flash Point

Value: 168 degree C Type: closed cup

Method: other: DIN 51758

Year:

Source: Bayer AG Leverkusen

16-SEP-1993 (2)

Value: 176.7 degree C

Type: open cup

Method: other: Cleveland Open Cup

Year:

26-SEP-2000 (7)

2.8 Auto Flammability

Value: 348.9 degree C

26-SEP-2000 (7)

2.9 Flammability

-

2.10 Explosive Properties

_

2.11 Oxidizing Properties

-

2.12 Additional Remarks

-

- 6/45 -

3. Environmental Fate and Pathways

3.1.1 Photodegradation

Type: air INDIRECT PHOTOLYSIS Sensitizer: OH

Conc. of sens.: 1560000 molecule/cm3

Rate constant: .0000000007949 cm3/(molecule * sec)

Degradation: 50 % after 1.6 hour(s)

Method: other (calculated): AOP Program vers1.89

1999 Year: GLP: no

Test substance: other TS: molecular structure Reliability: (2) valid with restrictions Accepted calculation method

Flaq: Critical study for SIDS endpoint

15-OCT-2001 (4)

Type: water Light source: Xenon lamp

DIRECT PHOTOLYSIS

Halflife t1/2: 26 minute(s)

Method:

Year: GLP:

Test substance:

Remark: light T1/2=26 minutes; Dark control T1/2=9.6 hours

Reliability: (2) valid with restrictions

15-OCT-2001 (3)

3.1.2 Stability in Water

abiotic Type:

Degradation: = 100 % after 24.9 hour(s)

at pH 7 and 20 degree C

Deg. Product: yes

Method: other: ABC Laboratory protocol; see TC Year: GLP: yes

Test substance: other TS: purity = 97% Remark: product = benzothiazole

Source: Monsanto

Bayer AG Leverkusen

Buffered deionized water; no light; initial concentra-Test condition:

tion = 1 mg/l

Reliability: (1) valid without restriction

GLP study; Meets generally accepted scientific method and is

described in sufficient detail

Flaq: Critical study for SIDS endpoint

15-OCT-2001 (8)

3.1.3 Stability in Soil

3.2 Monitoring Data (Environment)

- 7/45 -

Date: 17-OCT-2001
3. Environmental Fate and Pathways ID: 95-33-0

3.3.1 Transport between Environmental Compartments

Type: fugacity model level III

Media: other: air - biota - sediment(s) - soil - water

Air (Level I):
Water (Level I):
Soil (Level I):
Biota (L.II/III):
Soil (L.II/III):

Method: other: EPIWIN Level III Fugacity Model

Year: 1999

Result: Media Distribution Half-Life Emissions Fugacity $(percent) \qquad \qquad (hr) \qquad \qquad (kg/hr) \qquad (atm)$ 3.23 0.00923 Air 1000 2.33e-013 900 20.7 Water 1000 7.78e-015 1000 0 78.3 900 1.12e-014 Soil Sediment 0.924 3.6e+003 5.78e-015 0

> Persistence Time: 1.01e+003 hr Reaction Time: 1.27e+003 hr Advection Time: 4.8e+003 hr

Percent Reacted: 79
Percent Advected: 21

Reliability: (2) valid with restrictions
Accepted calculation method

Flag: Critical study for SIDS endpoint

15-OCT-2001 (4)

3.3.2 Distribution

-

3.4 Mode of Degradation in Actual Use

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- 8/45 -

Date: 17-OCT-2001 ID: 95-33-0

3. Environmental Fate and Pathways

3.5 Biodegradation

Type:

Inoculum: predominantly domestic sewage

Concentration: $100 \, \text{mg/l}$

Degradation: 0 % after 28 day

Result: under test conditions no biodegradation observed

Method: Directive 84/449/EEC, C.7 "Biotic degradation - modified MITI

test"

1988 GLP: no Year:

Test substance:

Remark: related to 02-consumption Bayer AG Leverkusen Source:

(1) valid without restriction Reliability: Meets National standards method

Critical study for SIDS endpoint Flaq:

15-OCT-2001 (2)

Type: aerobic

Inoculum: activated sludge, adapted

20 mg/l related to Test substance Concentration:

Degradation: ca. 0 % after 35 day

under test conditions no biodegradation observed Result: other: Method similar to ASTM draft method no. 2 ASTM Method:

Committee E35.24.

Year: 1979 GLP: no data

Test substance:

Remark: Method is similar in principle to Sturm test measuring

ultimate biodegradation as CO2 evolved

Source: Monsanto

Bayer AG Leverkusen

15-DEC-1995 (9)

Type: aerobic

Inoculum:

Degradation: 4 %

Method:

Year: GLP:

Test substance: other TS: purity = 97%

27-SEP-2000 (10)

3.6 BOD5, COD or BOD5/COD Ratio

Remark: ThOD: 2070 mg/g

Source: Bayer AG Leverkusen

05-OCT-1993 (2)

3.7 Bioaccumulation

- 9/45 -

Date: 17-OCT-2001
3. Environmental Fate and Pathways

ID: 95-33-0

3.8 Additional Remarks

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- 10/45 -

AQUATIC ORGANISMS

4.1 Acute/Prolonged Toxicity to Fish

Type: static

Species: Brachydanio rerio (Fish, fresh water)

Exposure period: 96 hour(s)

Unit: mg/l Analytical monitoring: no

LC0: >= 1000

Method: other: Letale Wirkung beim Zebrabaerbling,

UBA-Verfahrensvorschlag, Mai 1984, Letale Wirkung beim Zebrabaerbling Brachydanio rerio LCO, LC50, LC100, 48-96h

Year: 1989 GLP: no

Test substance:

Remark: direct weight

Source: Bayer AG Leverkusen

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

15-OCT-2001 (2)

Type: static

Species: Pimephales promelas (Fish, fresh water)

Exposure period: 96 hour(s)

Unit: mg/l Analytical monitoring: no

LC50: > 1000

Method: OECD Guide-line 203 "Fish, Acute Toxicity Test"
Year: 1984 GLP: yes

Test substance: other TS: purity = 97%

Remark: 24, 48, 96hr LC50 >1000 mg/l; water solubility=40 mg/l

Source: Monsanto

Bayer AG Leverkusen

Reliability: (1) valid without restriction

GLP guideline study

Flag: Critical study for SIDS endpoint

15-OCT-2001 (11)

Type: static

Species: Lepomis macrochirus (Fish, fresh water)

Exposure period: 96 hour(s)

Unit: mg/l Analytical monitoring: no

LC50: = 7.9

Method: other: Bionomics Lab protocol; see test conditions
Year: 1976 GLP: no data

Test substance:

Remark: C.I. for LC50=6.9-9.1 mg/l; 24 and 48hr LC50=8.8 mg/l

Source: Monsanto

Bayer AG Leverkusen

Test condition: Carrier-acetone; 22C; length=3.8cm; 15 L water; not fed;

no aeration

15-OCT-2001 (12)

- 11/45 -

Type: static

Species: Salmo gairdneri (Fish, estuary, fresh water)

Exposure period: 96 hour(s)

Unit: mg/l Analytical monitoring: no

LC50: = 5.4

Method: other: Bionomics Lab protocol; see test conditions
Year: 1976 GLP: no data

Test substance: other TS: purity 97%

Remark: C.I. for ELC50=4.5-6.5; 24hr LC50>7.5<10 mg/l; 48hr LC50=

7.5 mg/l

Source: Monsanto

Bayer AG Leverkusen

Test condition: Carrier-acetone; 15 L water; 10 fish/treatment; length=

2.7 cm; no feed; no aeration; 12 C

15-OCT-2001 (12)

Type: flow through

Species:

Exposure period: 14 day

Unit: mg/l Analytical monitoring: yes

Method: other: Springborn Lab protocol; see TC

Year: 1983 GLP: no data

Test substance:

Remark: 20 % mortality at 1.0 mg/l conc. (highest tested conc.)

Source: Monsanto

Bayer AG Leverkusen

Test condition: Nount-Brungs diluter; 19L aquaria; 10 fish/rep.; flow

rate=5 tanks volume/day; carrier-acetone; water solubility < 0.48 mg/l; two highest concentration above

water solubility

15-DEC-1995 (13)

Type: static

Species: Brachydanio rerio (Fish, fresh water)

Exposure period: 96 hour(s)

Unit: Analytical monitoring: yes

Method: other: Letale Wirkung beim Zebrabaerbling,

UBA-Verfahrensvorschlag, Mai 1984, Letale Wirkung beim Zebrabaerbling Brachydanio rerio LCO, LC50, LC100, 48-96h

Year: 1988 GLP: no

Test substance:

Remark: Analytical monitoring: DOC

Die Substanz wurde in der Konzentration 1000 mg/l 2h

eluiert, abfiltriert und das Filtrat getestet.

Keine Schadwirkung im Original.

DOC des Filtrats: 12 mg/l

Reliability: 2

Source: Bayer AG Leverkusen

27-SEP-2000 (2)

- 12/45 -

4.2 Acute Toxicity to Aquatic Invertebrates

Type:

Species: Daphnia magna (Crustacea)

Exposure period: 48 hour(s)

Unit: mg/l Analytical monitoring: no

NOEC: = 5.6 EC50: = 18

Method: OECD Guide-line 202, part 1 "Daphnia sp., Acute

Immobilisation Test"

Year: 1984 GLP: yes

Test substance:

Remark: C.I. for EC50=14-23 mg/l; 24hr EC50=21 mg/l

Source: Monsanto

Bayer AG Leverkusen

Reliability: (1) valid without restriction

GLP guideline study

Flag: Critical study for SIDS endpoint

15-OCT-2001 (14)

Type:

Species: Daphnia magna (Crustacea)

Exposure period: 96 hour(s)

Unit: mg/l Analytical monitoring: no

EC50: 18

Method:

Year: GLP: no data

Test substance: other TS: purity = 97%

27-SEP-2000 (15)

4.3 Toxicity to Aquatic Plants e.g. Algae

Species: Selenastrum capricornutum (Algae)

Endpoint: biomass
Exposure period: 96 hour(s)

Unit: mg/l Analytical monitoring: no

EC50: = .9 - 1.1

Method: OECD Guide-line 201 "Algae, Growth Inhibition Test" Year: 1984 GLP: yes

Test substance: other TS: purity = 97%

Remark: C.I. = 0.3-3.2 mg/l; in vivo chlorophyll: 24hr EC50= 3.2 mg/l, 48hr EC50=2.4 mg/l, 72hr EC50=1.2, 96hr EC50=

 $1.1~\mathrm{mg/l}$

Source: Monsanto

Bayer AG Leverkusen

Test condition: initial cell inoculum=20X3 cell/ml; 24C; 3800 lux;

carrier-DMF

Reliability: (1) valid without restriction

GLP quideline study

Flag: Critical study for SIDS endpoint

15-OCT-2001 (16)

- 13/45 -

4.4 Toxicity to Microorganisms e.g. Bacteria

Type: aquatic

Species: activated sludge

Exposure period: 3 hour(s)

Unit: mg/l Analytical monitoring: no

EC50: > 10000

Method: other: Test for Inhibition of Oxygen Consumption by Activated

Sludge, ISO 8192

Year: 1988 GLP: no

Test substance:

Remark: direct weight Reliability: 1

Source: Bayer AG Leverkusen

18-DEC-1995 (2)

Type: soil

Species:

Exposure period: 96 hour(s)

Unit: Analytical monitoring:

LC50 : Method:

Year: GLP:

Test substance:

Remark: test result: 25 %, when incorporated into nutrient

agar, moderately toxic

Source: Monsanto

Bayer AG Leverkusen

15-DEC-1995 (17)

4.5 Chronic Toxicity to Aquatic Organisms

4.5.1 Chronic Toxicity to Fish

-

4.5.2 Chronic Toxicity to Aquatic Invertebrates

-

- 14/45 -

TERRESTRIAL ORGANISMS

4.6.1 Toxicity to Soil Dwelling Organisms

-

4.6.2 Toxicity to Terrestrial Plants

-

4.6.3 Toxicity to other Non-Mamm. Terrestrial Species

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4.7 Biological Effects Monitoring

-

4.8 Biotransformation and Kinetics

_

4.9 Additional Remarks

_

- 15/45 -

5.1 Acute Toxicity

5.1.1 Acute Oral Toxicity

Type: LD50 Species: rat

Strain:

Sex: male/female

Number of

Animals: 5

Vehicle: other: corn oil Value: = 5300 mg/kg bw

Method: other

Year: 1973 GLP: no

Test substance: other TS: purity = 97%

Remark: test conditions:

mode of administration: gavage

vehicle: corn oil

number of animals: 5 per dose levels (2M, 3F, or 3M, 2F)

Result: Dercreased appetite and activity; weakness, tremors, collapse,

lung: hyperemia; liver: hyperemia;
gastro-intestinal tract: inflammation

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Flag: Critical study for SIDS endpoint

15-OCT-2001 (18) (19)

Type: LD50 Species: rat

Strain:
Sex:
Number of
Animals:
Vehicle:

Value: > 5000 mg/kg bw Method: other: no data

Year: 1977 GLP: no

Test substance: other TS: Soxinol CZ

Remark: 1000 mg/kg: No mortality and no toxic symptoms

2500 and 5000 mg/kg: 2/10 males and 3/10 females died 1-3 days after treatment with 5000 mg/kg. After 2-3 hours after

administration, irregular respiration, dyspnea,

hypersensitivity and ataxia were observed.

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Flag: Critical study for SIDS endpoint

15-OCT-2001 (20)

- 16/45 -

Type: LD50 Species: mouse

Strain:
Sex:
Number of
 Animals:
Vehicle:

Value: > 8000 mg/kg bw

Method: other

Year: GLP: no data

Test substance: other TS: Soxinol CZ-G
Remark: mortality: 0/8; only males
Reliability: (2) valid with restrictions

data are generally regarded as sufficient

Flag: Critical study for SIDS endpoint

15-OCT-2001 (21)

Type: LD50 Species: rat

Strain:
Sex:
Number of
Animals:
Vehicle:

Value: = 6850 mg/kg bw

Method: other

Year: GLP: no

Test substance: other TS: purity 96 to 98 %

Reliability: (3) invalid Reliability: 3

Remark: International Protocol Compliance: No

15-OCT-2001 (22)

Type: LD50 Species: mouse

Strain:
Sex:
Number of
Animals:
Vehicle:

Value: > 4000 mg/kg bw Method: other: no data

Year: GLP: no data

Test substance: other TS: no data Reliability: (4) not assignable

Reliability: 4

Remark: the test method is not stated

15-OCT-2001 (23)

5.1.2 Acute Inhalation Toxicity

-

5.1.3 Acute Dermal Toxicity

Type: LD50 Species: rabbit

Strain: Sex: Number of Animals:

Vehicle: other: corn oil Value: > 7940 mg/kg bw

Method: other

Year: 1973 GLP: no

Test substance: other TS: purity = 97%

Remark: mortality: 0/3 test conditions:

vehicle: corn oil (40 % suspension of test substance)

exposure time: 24 h

Result: no local effects noted; systemic: decreased appetite and

activity

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Flag: Critical study for SIDS endpoint

15-OCT-2001 (24) (19)

5.1.4 Acute Toxicity, other Routes

Type: LD50 Species: mouse

Strain:
Sex:
Number of
 Animals:
Vehicle:

Route of admin.: i.p.

Value: > 2500 mg/kg bw

Method: other

Year: GLP: no

Test substance: other TS: no data

Reliability: (2) valid with restrictions

Remark: data are generally regarded as sufficient

24-APR-2001 (25)

- 18/45 -

Type: LD50 Species: mouse

Strain:
Sex:
Number of
 Animals:
Vehicle:

Route of admin.: i.v.

Value: 32 mg/kg bw Method: other: no data

Year: GLP: no data

Test substance: other TS: no data
Reliability: (4) not assignable
Reliability: 4

Remark: data are derived from papers citing other papers

24-APR-2001 (26)

Type: LD50 Species: mouse

Strain:
Sex:
Number of
 Animals:
Vehicle:

Route of admin.: other: no data Value: > 8000 mg/kg bw

Method: other

Year: GLP: no data

Test substance: other TS: highest purity available in gum arabic

Reliability: (3) invalid Reliability: 3

Remark: documentation is not sufficient for an assessment

24-APR-2001 (27)

5.2 Corrosiveness and Irritation

5.2.1 Skin Irritation

Species: rabbit

Concentration:

Exposure:
Exposure Time:
Number of
Animals:
PDII:

Result: slightly irritating EC classificat.: not irritating

Method: other: Younger Laboratories, St Louis
Year: 1973 GLP: no data

Test substance: other TS: purity = 97%

Result: test result:

EU mean erythema score: 0.00 EU mean edema score: 0.00

- 19/45 -

PII: 0.00

other effects: none

healing time: not applicable

Test condition: test conditions:

mode of administration: not reported

exposure time: 24 hours number of animals: 6

Reliability: (2) valid with restrictions
Flag: Critical study for SIDS endpoint

15-OCT-2001 (18)

Species: rabbit

Concentration:

Exposure:
Exposure Time:
Number of
Animals:
PDII:

Result: slightly irritating

EC classificat.:

Method: other: Draize-Test

Year: GLP: yes

Test substance: other TS: as prescribed by chapter 1 in dataset of Monsanto

Result: test results:

EU mean erythema score: 0.17
EU mean edema score: 0.00

PII: 0.4/8

other effects: none healing time: 72 h

Test condition: test conditions:

mode of administration: occlusive

exposure time: 24 h number of animals: 6 skin: intact and abraded observation times: 24 h, 72 h

Reliability: (3) invalid

15-OCT-2001 (28) (19)

Species: rabbit

Concentration:

Exposure:
Exposure Time:
Number of
 Animals:
PDII:

Result: not irritating

EC classificat.:

Method: other: no data

Year: GLP: no data

Test substance: other TS: no data
Reliability: (4) not assignable

Reliability: 4

Remark: the test method is not stated

- 20/45 -

15-OCT-2001 (23)

Species: human

Concentration:

Exposure:
Exposure Time:
Number of
Animals:
PDII:
Result:

EC classificat.:

Method: other: no data

Year: GLP: no

Test substance: other TS: no data
Result: effect: no irritation
Reliability: (4) not assignable

Reliability: 4

Remark: the test method is not stated in detail

15-OCT-2001 (23)

5.2.2 Eye Irritation

Species: rabbit

Concentration:

Dose:

Exposure Time:
Comment:
Number of
Animals:

Result: slightly irritating

EC classificat.: not irritating

Method: OECD Guide-line 405 "Acute Eye Irritation/Corrosion"

Year: 1982 GLP: no

Test substance: other TS: purity = 97%

Result: test results:

EU mean erythema score: 1.7 EU mean chemosis score: 1.1

EU mean corneal opacity score: 0.0

EU mean iritis score: 0.0 Draize score: 1.8/110

other effects: discharge at 1 h and 24 h

Reliability: (1) valid without restriction

Guideline study

Flag: Critical study for SIDS endpoint

15-OCT-2001 (18) (19)

- 21/45 -

Species: rabbit

Concentration:

Dose:

Exposure Time:
Comment:
Number of
Animals:

Result: not irritating

EC classificat.:

Method: other: 5 mg/animal

Year: GLP: no

Test substance: other TS: no data Reliability: (4) not assignable

Reliability: 4

Remark: the test method is not stated in detail

24-APR-2001 (23)

5.3 Sensitization

Type: Buehler Test Species: guinea pig

Concentration: Induction 25 % open epicutaneous

Number of Animals: Vehicle:

Result: not sensitizing
Classification: not sensitizing
Method: other: see remarks

Year: GLP: yes

Test substance: other TS: Santocure accelerator

Method: induction exposure: a 25 % concentration of the test ma-

terial (in ethanol) was applied to the shaved skin of the test animals for 6 hours, once a week, for 3 consecutive weeks; challenge exposure: the test material was administered two weeks after the weekly applications were con-

cluded

Reliability: (1) valid without restriction

Remark: International Protocol Compliance: Yes

15-OCT-2001 (29)

Type: Patch-Test Species: human

Number of
Animals:
Vehicle:
Result:

Classification:

Method: other

Year: GLP: no data

Test substance: other TS: 1 % in petrolatum

Result: 34/686 patients had a positive patch test result with CBS

Reliability: (2) valid with restrictions

Remark: acceptable, well documented publication

24-APR-2001 (30)

- 22/45 -

Type: Patch-Test Species: human

Number of Animals: Vehicle: Result:

Classification:

Method: other

Year: GLP: no data

Test substance: other TS: 1 % in petrolatum

Result: 2/5 patients with contact dermatitis from rubber footwear had a positive patch test result with CBS among others

Reliability: (2) valid with restrictions

acceptable, well documented publication

24-APR-2001 (31)

Type: Patch-Test Species: human

Number of
Animals:
Vehicle:
Result:

Classification:

Method: other

Year: GLP: no data

Test substance: other TS: 1 % in petrolatum

Result: 11/46 patients with occupational rubber dermatitis had a

positive patch test result with CBS

Reliability: (2) valid with restrictions

acceptable, well documented publication

24-APR-2001 (32)

Type: Patch-Test Species: human

Number of
Animals:
Vehicle:
Result:

Classification:

Method: other

Year: GLP: no data

Test substance: other TS: 1 % in petrolatum

Result: 1/15 thiuram-sensitized patients had a positive patch test

result with CBS

Reliability: (2) valid with restrictions

acceptable, well documented publication

24-APR-2001 (33)

- 23/45 -

Type: Patch-Test Species: human

Number of
Animals:
Vehicle:
Result:

Classification:

Method: other: L. Schwartz and S. Peck, Public Health Reports, 59(17),

reprint 2552 (1944)

Year: 1944 GLP: no

Test substance: other TS: as prescribed by chapter 1 in dataset of Monsanto

Method: test conditions:

mode of application: occlusive, patch of 1.5 cm sq on upper

arm

exposure time: 24 hours for each exposure challenge: 10 days after first exposure

observation times: 24 and 48 hours after patch removal number of volunteers: 204 (94 F, 110 M), age ranging from

number of volunteers: 204 (94 F, 110 M), age ranging from 15 to 70 years $\,$

test results:

no positive reactions after first and second exposure

Reliability: (2) valid with restrictions

24-APR-2001 (34)

Type: Patch-Test Species: human

Number of Animals: Vehicle: Result:

Result:

Classification:

Method: other: L. Schwartz and S. Peck, Public Health Reports, 59(17),

reprint 2552 (1944)

Year: 1944 GLP: no

Test substance: other TS: as prescribed by chapter 1 in dataset of Monsanto

Result: test result:

no positive reactions on first and second exposure

Test condition: test conditions:

mode of administration: semi-occlusive(?), 1 inch sq patch

on upper arm

exposure time: 48 hours

observation times: 24, 48 and 72 hours after patch removal

challenge: 10 days after first exposure

vehicle: none

volunteers: 196 (100 M, 96 F), age ranging from 15 to 70

years

Reliability: (2) valid with restrictions

24-APR-2001 (35)

- 24/45 -

Type: Patch-Test Species: human

Number of Animals: Vehicle: Result:

Classification:

Method: other: Modified Shelanski

Year: GLP: no

Test substance: other TS: as prescribed by chapter 1 in dataset of Monsanto

Method: test conditions:

number of volunteers: 51, selected from local population mode of application: 0.2 g of material tested as a 70 % suspension in petrolatum was applied on the webril pad of a Parke-Davis Readi-bandage, the patch was then applied on the

skin (occlusive dressing)

induction: series of 12 applications of 24 h duration each

carried during weeks 1, 2 and 3.

challenge: a series of 4 applications on virgin sites at

week 6

Result: test results:

material acted as a sensitizer in 5 out of 51 volunteers

Reliability: (2) valid with restrictions

24-APR-2001 (36) (37)

Type: Patch-Test
Species: human

Number of
Animals:
Vehicle:
Result:

Classification:

Method:

Year: GLP: no data

Test substance: other TS: no data

Remark: Some patch tests carried out with the substance on people

suffering from contact dermatitis were positive

Reliability: (2) valid with restrictions

Remark: acceptable, well documented publication:

Bajaj, Foussereau, Rudzki

data are generally regarded as sufficient:

Kantoh, van Dijk, Holness

data are used as supporting evidence:

Eriksson, Heise

24-APR-2001 (38) (39) (40) (41) (42) (43) (44) (45)

- 25/45 -

Type: Patch-Test

Species: human

Number of
Animals:
Vehicle:
Result:

Classification:

Method:

Year: GLP: no data

Test substance: other TS: no data

Remark: One contact dermatitis patient had a negative patch test

result with CBS.

Reliability: (2) valid with restrictions

Remark: acceptable, well documented publication

24-APR-2001 (46)

Type: other Species: human

Number of
Animals:
Vehicle:
Result:

Classification:

Method:

Year: GLP:

Test substance: other TS: CBS

Result: There are numerous reports in the literature of skin

sensitization to CBS; however, in most cases the patients showed positive reactions to other constituents of the "mercapto mix" used for skin testing

(MBT, MBTS and MDR)

Reliability: (4) not assignable

Remark: the data quality and test result cannot be evaluated

24-APR-2001 (47) (48) (49) (50) (51) (52)

Type: other: closed epicutaneous test

Species: guinea pig

Number of
 Animals:
Vehicle:

Result: not sensitizing

Classification:

Method: other: induction: 0,5 M (50 mg), challenge: 0,5 M or 0,05 M

Year: GLP: no data

Test substance: other TS: no data

Reliability: (2) valid with restrictions

Remark: data are generally regarded as sufficient

24-APR-2001 (43)

- 26/45 -

5.4 Repeated Dose Toxicity

Species: rat Sex: male/female

Strain: no data Route of admin.: oral feed

Exposure period: 4 w

Frequency of

treatment: daily

Post. obs.

period: no data

Doses: 100, 250, 500, 1000 or 3000 mg/kg bw/d

Control Group: yes

NOAEL: = 250 mg/kg bw

Method:

Year: GLP: yes

Test substance: other TS: Santocure accelerator

Result: evidence of toxicity, as indicated by reduced body weight

gains and food consumption, was noted at the 500, 1000,

and 3000 mg/kg bw/d exposure levels

Reliability: (2) valid with restrictions

Remark: No hematology, no blood biochemistry, no

histopathology

Flag: Critical study for SIDS endpoint

15-OCT-2001 (53)

Species: rat Sex: no data

Strain: no data
Route of admin.: inhalation

Exposure period: 4 w

Frequency of

treatment: 6 h/d, 5 d/w

Post. obs.

period: no data

Doses: 0.0043, 0.0144 or 0.048 mg/l

Control Group: yes

NOAEL: = .0144 mg/l

Method:

Year: GLP: yes

Test substance: other TS: Santocure accelerator

Result: elevated clinical chemistry (SGOT) values were observed

in mid- and high-exposure animals; microscopic lesions in the conjunctiva, lymph nodes and spleen were noted for high-exposure group animals at an incidence greater

than control animals

Reliability: (1) valid without restriction

International Protocol Compliance: Yes

Flag: Critical study for SIDS endpoint

15-OCT-2001 (54)

- 27/45 -

Species: rabbit Sex: no data

Strain: no data Route of admin.: dermal Exposure period: 21 d

Frequency of

treatment: daily

Post. obs.

period: no data

Doses: 125, 500 or 2000 mg/kg bw/d

Control Group: yes

Method:

Year: GLP: yes

Test substance: other TS: Santocure accelerator

Result: no evidence of toxicity related to test material admin-

istration

Reliability: (1) valid without restriction

International Protocol Compliance: Yes

Flag: Critical study for SIDS endpoint

15-OCT-2001 (55)

Species: rat Sex: no data

Strain: no data
Route of admin.: inhalation
Exposure period: 15 days

Frequency of

treatment: 24h/day, daily

Post. obs.

period: no

Doses: 300-400 mg/m3
Control Group: no data specified

Method:

Year: GLP: no data

Test substance: other TS: no data

Result: body weight gain unchanged, in few animals slight transient

changes in the functional state of the nervous system

Reliability: (4) not assignable

Remark: the test method is not stated in detail

24-APR-2001 (23)

- 28/45 -

Species: rat Sex: no data

Strain: no data Route of admin.: gavage

Exposure period: 24 d during a period of 5 w

Frequency of

treatment: daily

Post. obs.

period: no data

Doses: 0.5 or 1.25 mg/kg bw/d

Control Group: yes

Method:

Year: GLP: no

Test substance: other TS: Santocure accelerator

Result: two females in the high-dose group and one female in

the low-dose group died during the experiment; a decrease in body weight gain in the treated animals and an increase in relative thyroid weights were observed; no pathologic changes attributable to treat-

ment were noted

Reliability: (3) invalid

International protocol compliance: no

24-APR-2001 (22)

Species: rat Sex: male/female

Strain: other: H.L.A.
Route of admin.: oral unspecified

Exposure period: 5w

Frequency of

treatment: 5d/w, daily

Post. obs.

period: 4d

Doses: 500 or 1250 mg/kg bw

Control Group: other: yes

Method: other

Year: GLP: no

Test substance: other TS: purity 96 to 98 %

Result: 1250 mg/kg (females): increased mortality,

hyperirritability, sanguineous discharge from the mouth and

nostrils; females of both dose groups: changes of the thyroid weight, without pathologic changes (microscopic

examination)

Reliability: (3) invalid

International protocol compliance: no

24-APR-2001 (22)

- 29/45 -

Species: mouse Sex: male/female

Strain: other: ddy/Slc Route of admin.: oral feed Exposure period: 3 months

Frequency of

treatment: daily

Post. obs.

period: no

Doses: 0, 0.04, 0.2, 1.0 %

Control Group: yes, concurrent no treatment

Method: other

Year: GLP: no data

Test substance: other TS: Soxinol CZ-G

Remark: English Translation of an Informal Japanese report with

figures and tables

Result: no mortality is reported; reduced body weight gain (1%-dose

group); no dose despended changes of hematological/

biochemical parameters and organ weights; no

histopathological examinations reported

Reliability: (3) invalid

Remark: documentation is not sufficient for an assessment

24-APR-2001 (21)

Species: rabbit Sex: no data

Strain: no data

Route of admin.: oral unspecified

Exposure period: 3,5 months

Frequency of

treatment: every second day for the first 1,5 months followed by daily

dosing

Post. obs.

period: no

Doses: 20 mg/kg bw Control Group: other: yes

Method:

Year: GLP:

Test substance:

Result: no particular symptoms Reliability: (4) not assignable

Remark: the test method is not stated in detail

24-APR-2001 (23)

- 30/45 -

5.5 Genetic Toxicity 'in Vitro'

Ames test Type:

System of

testing: no data

Concentration: Cytotoxic Conc.:

Metabolic

with and without activation:

negative Result: Method: other:

Year: GLP: no

Test substance: other TS: Santocure accelerator Reliability: (2) valid with restrictions

Remark: International Protocol Compliance: +/- Yes

Flaq: Critical study for SIDS endpoint

17-OCT-2001 (56)

Type: Mouse lymphoma assay

System of

testing: L5178Y mouse lymphoma assay Concentration: 0.078 ug/ml to 40 ug/ml

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method: other: Litton Bionetics method

Year: 1979 GLP: no

Test substance: other TS: purity = 97%

Remark:

test conditions:

solubility in medium: precipitation at 78 ug/ml and higher

cytotoxicity:

-S9:

no survivors at 39 ug/ml, 5.7% to 13.3% relative growth for

the concentration range from 15 to 30 ug/ml

no survivors at 80 ug/ml, 75.2% relative growth at 40 ug/ml,

toxicity was moderate to high in 0.313 to 20 ug/ml

concentration range

(2) valid with restrictions Reliability:

Flag: Critical study for SIDS endpoint

15-OCT-2001 (57)

- 31/45 -

Type: Ames test

System of

testing: Salmonella typhimurium, TA 1535, TA 1537, TA 1538, TA 98, TA

100

Concentration: 0.1 - 500 ug/plate

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method: other: no data

Year: GLP: no

Test substance: other TS: no data

Reliability: (3) invalid

Remark: data are derived from reviews

15-OCT-2001 (58)

Type: Ames test

System of

testing: no data Concentration: no data

Cytotoxic Conc.:

Metabolic

activation: no data Result: negative

Method: other: no data

Year: GLP: no data

Test substance: other TS: no data Reliability: (4) not assignable

Remark: data are derived from papers citing other papers

see Rannug et al. (1984)

24-APR-2001 (59)

Type: Ames test

System of

testing: TA 100, TA 98

Concentration: no data

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method: other: no data

Year: GLP: no data

Test substance: other TS: no data Reliability: (4) not assignable

Remark: data are derived form abstracts

24-APR-2001 (60)

- 32/45 -

Type: Ames test

System of

testing: Salmonella typhimurium TA 98, TA 100, TA 1535, TA 1537, TA

1538

Concentration: 10-3000 ug/plate

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method: other: no data

Year: 1981 GLP: no data

Test substance: other TS: Soxinol CZ Reliability: (4) not assignable

Remark: the data quality and test result cannot be evaluated

24-APR-2001 (61)

Type: Ames test

System of

testing: Salmonella typhimurium, TA 1535, TA 1537, TA 1538, TA 98, TA

100

Concentration: 0 - 150 ug/plate (-S9); 0 - 200 ug/plate (+S9)

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method: other: no data

Year: GLP: no data

Test substance: other TS: technical purity Reliability: (2) valid with restrictions

Remark: acceptable, well documented publication

24-APR-2001 (62)

Type: Yeast gene mutation assay

System of

testing: Saccharomyces cerevisiae

Concentration: 0.1 - 500 ug/plate

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method: other: no data

Year: GLP: no

Test substance: other TS: no data

Reliability: (3) invalid

Remark: data are derived from reviews

24-APR-2001 (58)

- 33/45 -

5.6 Genetic Toxicity 'in Vivo'

Type: other

Species: rat Sex: male/female

Strain:

Route of admin.: oral unspecified

Exposure period: twice (1st, 3rd day of oestrus)

Doses: 2000 mg/kg bw

Result:

Method: other

Year: GLP: no data

Test substance: other TS: Santocure

Remark: incomplete and inexact data on study design

Result: No increased postimplantation embryonic mortality (= "index

of mutagenicity").

Reliability: (3) invalid

Remark: unsuitable test system

24-APR-2001 (63)

5.7 Carcinogenicity

Species: mouse Sex: male/female

Strain: other: B6C3F1 and B6AKF1

Route of admin.: oral unspecified

Exposure period: 18 months

Frequency of

treatment: daily

Post. obs.

period: no

Doses: see remarks
Result: negative
Control Group: other: yes

Method: other

Year: GLP: no

Test substance: other TS: Durax

Remark: dose: 7th to 28th day of age: 215 mg/kg bw/d by stomach

tube, thereafter administration of 692 mg/kg diet

(app.: 99 mg/kg bw/d)

Result: no increased tumor incidence Reliability: (2) valid with restrictions

Remark: deviations from standard guidelines are acceptable

with regard to the creation date of the study

17-OCT-2001 (64) (65)

- 34/45 -

Species: mouse Sex: male/female

Strain: other: B6C3F1 and B6AKF1

Route of admin.: s.c. Exposure period: 1 d

Frequency of

treatment: once

Post. obs.

period: 18 months

Doses: 1000 mg/kg bw

Result: negative

Control Group: other: yes

Method: other

Year: GLP: no

Test substance: other TS: Durax

Result: no increased tumor incidence Reliability: (2) valid with restrictions

Remark: deviations from standard guidelines are acceptable

with regard to the creation date of the study

17-OCT-2001 (64)

5.8 Toxicity to Reproduction

Type: Fertility

Species: rat Sex: female

Strain:

Route of admin.: oral feed

Exposure Period: day 0-20 of gestation

Frequency of

treatment: daily

Duration of test:

Doses: 0, 0.001%, 0.01%, 0.1% and 0.5%.

Control Group: yes
NOAEL Parental: .01 %
NOAEL F1 Offspr.: .1 %

Method:

Year: GLP:

Test substance:

Result: Average daily intakes were 0, 0.7, 7.1, 69.6 and 288.8 mg/kg

respectively. Significantly lower maternal body weight gain, food consumption and fetal body weight means were noted in the 0.5% group. Maternal body weight gains were significantly reduced in the 0.1% group. There were NO

significant compound-related effects on pre- and

post-implantation losses, the number and ratio of live to

dead fetuses, or terata.

Flag: Critical study for SIDS endpoint

24-APR-2001 (66)

- 35/45 -

Type: other

Species: rat Sex: female

Strain: no data

Route of admin.: oral unspecified

Exposure Period: daily

Frequency of

treatment: 1st to 3rd day of estrus

Duration of test:

Doses: 2000 mg/kg bw/d

Control Group: yes

NOAEL Parental: 2000 mg/kg bw

Method: other

Year: GLP: no data

Test substance: other TS: Santocur

Remark: post observation period: up to the 19th day of pregnancy lowered weight of fetuses, no change in fertility, no

visible signs of poisoning in dams

Reliability: (3) invalid

Remark: unsuitable test system

24-APR-2001 (63)

5.9 Developmental Toxicity/Teratogenicity

Species: rat Sex: female

Strain: other: Charles river COBS CD

Route of admin.: gavage

Exposure period: 6th to 15th day of pregnancy

Frequency of

treatment: daily

Duration of test:

Doses: 100, 300, 500, 900 mg/kg bw/d

Control Group: yes

NOAEL Maternalt.: = 100 mg/kg bw NOAEL Teratogen.: = 500 mg/kg bw

Method: OECD Guide-line 414 "Teratogenicity"
Year: 1981 GLP: yes

Test substance: other TS: N-cyclohexyl-2-benzothiazole; purity = 97%

Remark: 1) 100 mg/kg bw:

NOEL

2) 300 mg/kg bw:

maternal: body weight gain, decr.

3) 500 mg/kg bw:

maternal: alopecia; body weight gain, decr.;

body weight, decr.

offspring: fetus, weight, decr.;

NOEl, teratogenic effects

4) 900 mg/kg bw:

maternal: excessive toxicity; death

Reliability: (1) valid without restriction

GLP Guideline study

Flag: Critical study for SIDS endpoint

17-OCT-2001 (67)

- 36/45 -

Species: rat Sex: female
Strain: other: Institute's own breeding colony (Imp: DAK)

Route of admin.: gavage

Exposure period: days 6 to 15 of gestation

Frequency of

treatment: once a day

Duration of test: day 20 of gestation

Doses: 50, 150 and 450 mg/kg

Control Group: yes, concurrent vehicle

NOAEL Maternalt.: 150 mg/kg bw NOAEL Teratogen.: 50 mg/kg bw

Method: other

Year: GLP: no data

Test substance:

Result: 450 mg/kg: decreased body weight gain, increased relative

kidney weights, decreased absolute spleen weights of the dams; 150 and 450 mg/kg: dose dependent increased number of fetuses with internal hydrocephalus; the number of fetuses with enlarged cerebral ventricles and/or renal pelvis was

not dose dependendent increased

24-APR-2001 (68)

Species: rat Sex: female

Strain: Wistar Route of admin.: oral feed

Exposure period: day 0 to day 20 of pregnancy

Frequency of

treatment: daily

Duration of test:

Doses: 0.7; 7.1; 69.6 or 288.8 mg/kg bw/d

Control Group: yes

NOAEL Maternalt.: 69.6 mg/kg bw NOAEL Teratogen.: 288.8 mg/kg bw

Method: other

Year: GLP: no data

Test substance: other TS: Soxinol CZ-G, 99 % pure

Remark: dose: 0.001; 0.01; 0.1 or 0.5 % in the diet

Result: except reduced food consumption, lowered weight of fetuses

and of the placentae in the highest dosage group no sings of toxicity in any group; no embryotoxic or teratogenic

effects

Reliability: (2) valid with restrictions

Remark: acceptable, well documented publication

24-APR-2001 (69)

- 37/45 -

Species: rat Sex: female

Strain:

Route of admin.: oral feed

Exposure period: days 6-15 of gestation

Frequency of treatment:
Duration of test:

Doses: 0, 100, 300, or 500 mg/kg

Control Group: yes

NOAEL Maternalt.: 300 mg/kg bw NOAEL Teratogen.: 300 mg/kg bw

Method:

Year: GLP:

Test substance: other TS: N-cyclohexyl-2-benzothiazole; purity = 97%

Result: Fetal body weight means and maternal body weight gains were

significantly reduced in the high dosage groups. Maternal general toxicity: None below 300 mg/kg

Foetal data: No effects below 300 mg/kg

24-APR-2001 (70)

Species: rat Sex: female

Strain: no data

Route of admin.: oral unspecified

Exposure period: daily

Frequency of

treatment: 4th to 11th day of pregnancy

Duration of test:

Doses: 2000 mg/kg bw/d Control Group: other: yes

Method: other

Year: GLP: no data

Test substance: other TS: Santocur

Remark: post exposure period: up to the 19th day of pregnancy lowered weight of fetuses, increased embryonic mortality,

no visible signs of poisoning in dams, increased

postimplantation embryonic mortality

Reliability: (3) invalid

Remark: unsuitable test system

24-APR-2001 (63)

- 38/45 -

Species: other Sex: no data

Strain:

Route of admin.:
Exposure period:
Frequency of
 treatment:
Duration of test:

Doses:

Control Group:

Method:

Year: GLP:

Test substance:

Remark: The substance was injected into the heart of 3-day chicken

embryos

Result: some malformations

Reliability: (3) invalid

Remark: test method not validated

24-APR-2001 (71)

Species: other Sex: no data

Strain:

Route of admin.:
Exposure period:
Frequency of
 treatment:
Duration of test:

Doses:

Control Group:

Method:

Year: GLP:

Test substance:

Remark: 3-day old chicken embryos received a maximum dose of

1 mole/egg (injected into the air-chamber); the relevance of this test system in relation to mammals

remains questionable

Result: 13 % malformed embryos

Reliability: (3) invalid

Remark: test method not validated

24-APR-2001 (72)

5.10 Other Relevant Information

Type: Metabolism

Remark: About 92% of the administred radioactivity was recovered

from urin and feces within 3 days after administration of 14C-N-cyclohexyl-2-benzothiazol-sulfenamide. No specific organ affinity was observed. Metabolites were cyclo-

hexylamine and 2-mercaptobenzothiazole.

Reliablity: 2

Remark: data are generally regarded as sufficient

Source: Bayer AG Leverkusen

24-NOV-1995 (73)

- 39/45 -

Type: other

Remark: in a 4-week comedogenicity (acnegenicity) assay,

Santocure vulcanization accelerator was applied to the inner surface of rabbit ears 5 days per week at concentrations of 0.01, 0.1 or 10 % in chloroform; no production of comedones was observed after repea-

ted treatment

Reliability: 3 GLP: No

Remark: International Protocol Compliance: No

Source: Monsanto

Bayer AG Leverkusen

23-APR-1996 (74)

Type: other

Remark: Reliability: 2

Remark: acceptable, well documented publication

when MBT, MBTS, MMBT and CBS (= mercapto mix patch testing standard) were dissolved in petrolatum or in buffer of

physilological pH, they were mainly converted into the redox

pair MBT and MBTS

Source: Bayer AG Leverkusen

24-NOV-1995 (75)

Type: other: metabolism/pharmacokinetics

Remark: A single oral dose of 250 mg 14C-CBS/rat (3m):

ca. 47% radioactivity in urine and 45 % in faeces

(total recovery after 3 days=92%)

2h after administration no other metabolite than MBTS (65%)

was observed in the stomach

Incubation with artifical gastric juice:

Transformation to MBT and MBTS

Reliability: 3

Remark: poor or deficient study design

Source: Bayer AG Leverkusen

25-APR-1996 (76)

5.11 Experience with Human Exposure

Remark: Occupational exposure to Santocure has been reported

to cause irritation to the eyes, skin and upper re-

spiratory tract
Reliability: 2

Source: Monsanto

Bayer AG Leverkusen

26-APR-1996 (77)

- 40/45 -

Date: 17-OCT-2001
6. References ID: 95-33-0

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Date: 17-OCT-2001 6. References ID: 95-33-0

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Date: 17-OCT-2001
6. References ID: 95-33-0

6. References ID: 95-33-0

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Date: 17-OCT-2001 6. References ID: 95-33-0

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- 44/45 -

7. Risk Assessment Date: 17-OCT-2001 ID: 95-33-0

7.1 End Point Summary

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7.2 Hazard Summary

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7.3 Risk Assessment

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- 45/45 -

I U C L I D

Data Set

Existing Chemical ID: 102-77-2 CAS No. 102-77-2

EINECS Name 2-(morpholinothio)benzothiazole

EINECS No. 203-052-4

TSCA Name Morpholine, 4-(2-benzothiazolylthio)-

Molecular Formula C11H12N2OS2

Producer Related Part

Company:

Creation date: 15-JUL-1999

Substance Related Part

Company:

Creation date: 15-JUL-1999

Memo: Rubber and Plastic Additives (RAPA) HPV Panel

Printing date: 15-OCT-2001

Revision date:

Date of last Update: 15-OCT-2001

Number of Pages: 60

Chapter (profile): Chapter: 1, 2, 3, 4, 5, 7

Reliability (profile): Reliability: without reliability, 1, 2, 3, 4

Flags (profile): Flags: without flag, confidential, non confidential, WGK

(DE), TA-Luft (DE), Material Safety Dataset, Risk

Assessment, Directive 67/548/EEC, SIDS

Date: 15-OCT-2001

1. General Information ID: 102-77-2

1.0.1 OECD and Company Information

Type: lead organisation

Name: American Chemistry Council (formerly Chemical Manufacturers

Association) Rubber and Plastic Additives (RAPA) HPV Panel

Street: 1300 Wilson Boulevard Town: 22209 Arlington, VA

Country: United States
Phone: 703-741-5600
Telefax: 703-741-6091

12-OCT-2001

Type: cooperating company
Name: Bayer Corporation
Country: United States

12-OCT-2001

Type: cooperating company

Name: Ciba Specialty Chemicals Corporation

Country: United States

12-OCT-2001

Type: cooperating company Name: Crompton Corporation

Country: United States

12-OCT-2001

Type: cooperating company Name: Flexsys America L.P.

Country: United States

12-OCT-2001

Type: cooperating company

Name: Noveon, Inc (formerly BF Goodrich)

Country: United States

12-OCT-2001

Type: cooperating company

Name: R.T. Vanderbilt Company, Inc.

Country: United States

12-OCT-2001

Type: cooperating company

Name: The Goodyear Tire & Rubber Company

Country: United States

12-OCT-2001

- 1/60 -

Date: 15-OCT-2001

1. General Information

Date: 15-OCT-2001

1D: 102-77-2

Type: cooperating company
Name: The Lubrizol Corporation

Country: United States

12-OCT-2001

Type: cooperating company

Name: UOP, LLC. Country: United States

12-OCT-2001

1.0.2 Location of Production Site

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1.0.3 Identity of Recipients

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1.1 General Substance Information

Substance type: organic
Physical status: solid
Purity: > 93 % w/w

20-OCT-1999

1.1.0 Details on Template

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1.1.1 Spectra

-

1.2 Synonyms

2-morpholinothio-benzothiazole 20-OCT-1999

N-oxydiethylene-2-benzothiazolesulfenamide 20-OCT-1999

- 2/60 -

Date: 15-OCT-2001

1. General Information

Date: 15-OCT-2001

10: 102-77-2

1.3 Impurities

CAS-No: 110-91-8
EINECS-No: 203-815-1
EINECS-Name: morpholine
Contents: < .4 % w/w

Remark: others: disulphides and sulfonic acid derivatives of

mercaptobenzothiazole, dimercaptobenzothiazole, and

methylmercaptobenzothiazole < 6%</pre>

20-OCT-1999

1.4 Additives

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1.5 Quantity

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1.6.1 Labelling

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1.6.2 Classification

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1.7 Use Pattern

Type: type

Category: Use resulting in inclusion into or onto matrix

20-OCT-1999

Type: industrial

Category: Polymers industry

20-OCT-1999

Type: use

Category: Vulcanizing agents

20-OCT-1999

1.7.1 Technology Production/Use

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1.8 Occupational Exposure Limit Values

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1.9 Source of Exposure

_

- 3/60 -

Date: 15-OCT-2001

1. General Information

Date: 15-OCT-2001

1D: 102-77-2

- 1.10.1 Recommendations/Precautionary Measures
- 1.10.2 Emergency Measures

1.11 Packaging

-

1.12 Possib. of Rendering Subst. Harmless

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1.13 Statements Concerning Waste

_

1.14.1 Water Pollution

_

1.14.2 Major Accident Hazards

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1.14.3 Air Pollution

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1.15 Additional Remarks

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1.16 Last Literature Search

-

1.17 Reviews

_

1.18 Listings e.g. Chemical Inventories

-

- 4/60 -

2.1 Melting Point

Value: 150.7 degree C

Method: other: MPBPWIN (v1.31)

Year: 1999 GLP: no

Testsubstance: other TS: molecular structure

Melting Point: 308.70 deg C (Adapted Joback Method) Result: Melting Point: 111.17 deg C (Gold and Ogle Method)

Mean Melt Pt : 209.94 deg C (Joback; Gold,Ogle Methods)

Selected MP: 150.68 deg C (Weighted Value)

Reliability: (2) valid with restrictions Accepted calculation method

Critical study for SIDS endpoint Flaq:

15-OCT-2001 (1)

75 - 90 degree C Value:

29-SEP-2000 (2)

Value: > 78 degree C

24-APR-2001 (3)

79 degree C Value:

29-SEP-2000 (4)

2.2 Boiling Point

Value: 385.1 degree C

Method: other: MPBPWIN (v1.31) Adapted Stein and Brown Method

Year: 1999 GLP: no

Testsubstance: other TS: molecular structure (2) valid with restrictions Reliability: Accepted calculation method

Flaq: Critical study for SIDS endpoint

15-OCT-2001 (1)

Value:

Decomposition: yes

Critical study for SIDS endpoint Flag:

15-OCT-2001 (3)

2.3 Density

relative density Type:

Value: 1.35 g/cm3 at 20 degree C

24-APR-2001 (3)(4)

2.3.1 Granulometry

- 5/60 -

2. Physico-chemical Data

2.4 Vapour Pressure

Value: .000001346 hPa at 25 degree C Method: other (calculated): MPBPWIN (v1.31)

Year: 1999 GLP: no

other TS: molecular structure Testsubstance:

Result: Vapor Pressure Estimations (25 deg C): (Using BP: 385.05 deg C (estimated))

(Using MP: 150.68 deg C (estimated)) VP: 2.79E-007 mm Hg (Antoine Method)

VP: 1.01E-006 mm Hg (Modified Grain Method)

VP: 2.22E-006 mm Hg (Mackay Method)

Selected VP: 1.01E-006 mm Hg (Modified Grain Method)

Reliability: (2) valid with restrictions Accepted calculation method

Flag: Critical study for SIDS endpoint

15-OCT-2001 (1)

Value: .0000023 hPa at 20 degree C

24-APR-2001 (3)

.0000045 hPa at 25 degree C Value:

24-APR-2001 (3)

2.5 Partition Coefficient

log Pow: 1.025 at 25 degree C

Method: other (calculated): KOWWIN Program (v1.65)

Year: 1999 GLP: no

Testsubstance: other TS: molecular structure (2) valid with restrictions Reliability: Accepted calculation method

Critical study for SIDS endpoint Flaq:

15-OCT-2001 (1)

log Pow: 3.49

Method: Year:

Testsubstance: other TS: Santocure MOR Accelertor

Remark: Pow: 3100

Flag: Critical study for SIDS endpoint

15-OCT-2001 (5)

- 6/60 -

Date: 15-OCT-2001 ID: 102-77-2 2. Physico-chemical Data

2.6.1 Water Solubility

Value: 3061 mg/l at 25 degree C Method: other: WSKOW (v1.36)

1999 Year: no GLP:

Testsubstance: other TS: molecular structure Reliability: (2) valid with restrictions
Accepted calculation method

Critical study for SIDS endpoint Flag:

(1)15-OCT-2001

Value: 32 other: pp Qualitative: not soluble 32 other: ppm

Testsubstance: other TS: Sanatocure MOR Accelerator

15-OCT-2001 (6)

2.6.2 Surface Tension

2.7 Flash Point

Value: 188 degree C
Type: open cup
Method: other: DIN 53

other: DIN 51584 Method:

Year:

24-APR-2001 (3)

2.8 Auto Flammability

2.9 Flammability

2.10 Explosive Properties

2.11 Oxidizing Properties

2.12 Additional Remarks

- 7/60 -

Date: 15-OCT-2001 ID: 102-77-2 3. Environmental Fate and Pathways

3.1.1 Photodegradation

Type: air INDIRECT PHOTOLYSIS Sensitizer: OH

Conc. of sens.: 156000 molecule/cm3

Rate constant: .000000001199482 cm3/(molecule * sec)

Degradation: 50 % after 1.1 hour(s)

Deg. Product: not measured

Method: other (calculated): AOP Program (v1.89)

Year: 1999 GI.P: no

Test substance: other TS: molecular structure Reliability: (2) valid with restrictions Accepted calculation method

Flaq: Critical study for SIDS endpoint

15-OCT-2001 (1)

Type: water Sun light Light source:

Spectr.of subst.: lambda (max, >295nm): 300 nm Conc. of subst.: 1.006 mg/l at 26 degree C

DIRECT PHOTOLYSIS

Halflife t1/2: 1 hour(s)

Degradation: 97 % after 4 hour(s)

Method: other (measured): Method similar to ASTM draft method No. 6,

ASTM E35.24 Subcommittee Aqueous Photolysis Task Group

1980 Year: GLP: no data

Test substance: other TS: Santocure MOR

Remark: Dark control recovery was 97 % at 4 hours while Santocurs

MOR values at 0, 1, 2, 3 and 4 hours were 100 %, 45 %, 15 %,

8 % and 3 %, respectively.

(2) valid with restrictions Reliability: Flag: Critical study for SIDS endpoint

24-APR-2001 (7)

- 8/60 -

3.1.2 Stability in Water

abiotic Type:

Degradation: 24 % after 25 hour(s)

at pH 7

Method: other: Hydrolysis in pH buffered dionized water

1984 Year: GLP: yes

Test substance: other TS: Santocure MOR

Remark: Primary purpose of this study was to identify hydrolysis by

> products. After 7 days of exposure to pH 7 water the following by products were determined to be present in the percentages listed in parenthesis: benzothiazole (64 %), mercaptobenzothiazole (21 %), unknown with proposed

composition of C11H14S2N2O2 (15 %) proposed structure shown below and morpholine (% not listed). Hydrolysis was complete

after 7 days at pH 7.

Reliability: (1) valid without restriction

GLP study; Meets generally accepted scientific method and is

described in sufficient detail

Flaq: Critical study for SIDS endpoint

15-OCT-2001 (8)

3.1.3 Stability in Soil

3.2 Monitoring Data (Environment)

3.3.1 Transport between Environmental Compartments

Type: fugacity model level III

Media: other: air, watr, soil, sediment

Air (Level I): Water (Level I): Soil (Level I): Biota (L.II/III): Soil (L.II/III):

Reliability:

other: (calculation) EPIWIN Level III Fugacity Model Method:

Year: 1999

Result: Media Distribution Half-Life Emissions Fugacity

> (percent) (hr) (kg/hr) (atm) 9.47e-005 2.14 1000 2.15e-015 Air Water 44.6 900 1000 1.19e-016 1000 55.3 4.06e-015 Soil 900 Sediment 0.0904 1.09e-016 3.6e+003 0

Persistence Time: 823 hr Reaction Time: 1.3e+003 hr Advection Time: 2.24e+003 hr

Percent Reacted: 63.3 Percent Advected: 36.7 (2) valid with restrictions

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3. Environmental Fate and Pathways

Flaq: Critical study for SIDS endpoint

15-OCT-2001 (1)

3.3.2 Distribution

3.4 Mode of Degradation in Actual Use

3.5 Biodegradation

Type: aerobic

Inoculum: predominantly domestic sewage

Concentration: 100 mg/l related Degradation: 0 % after 28 day 100 mg/l related to COD (Chemical Oxygen Demand)

Method: Directive 84/449/EEC, C.7 "Biotic degradation - modified MITI

test"

1988 Year: GLP: no

Test substance: other TS: 2-morpholinothio-benzothiazole; purity > 93%

Reliability: (1) valid without restriction Meets National standards method

Critical study for SIDS endpoint Flag:

15-OCT-2001 (3)

3.6 BOD5, COD or BOD5/COD Ratio

ThOD: 1880 mg/g Remark:

24-APR-2001 (3)

3.7 Bioaccumulation

Species: other

Exposure period: Concentration:

BCF: 1.23

Elimination:

Method: other: BCF Program (v2.13)

1999 Year: GLP: no

other TS: molecular structure Test substance: Log Kow (estimated) : 1.02Result:

Log Kow (experimental): not available from database

Log Kow used by BCF estimates: 1.02

Equation Used to Make BCF estimate: Log BCF = 0.77 log Kow - 0.70

Estimated Log BCF = 0.089 (BCF = 1.227)

(2) valid with restrictions Reliability:

Accepted calculation method

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Date: 15-OCT-2001
3. Environmental Fate and Pathways

ID: 102-77-2

15-OCT-2001 (1)

3.8 Additional Remarks

-

- 11/60 -

AQUATIC ORGANISMS

4.1 Acute/Prolonged Toxicity to Fish

Type: static

Species: Pimephales promelas (Fish, fresh water)

Exposure period: 96 hour(s)

mg/1Analytical monitoring: no

NOEC: = 1 LC50: = 3.5

OECD Guide-line 203 "Fish, Acute Toxicity Test" Method: Year: 1984

Test substance: other TS: Santocure MOR

Remark: C.I. for 96 h LC50 = 2.7-4.4 mg/l; 24 and 48 h LC50 = 4.0

Test condition: length = 25.8 mm; weight = 0.26 gReliability: (1) valid without restriction

GLP guideline study

Flag: Critical study for SIDS endpoint

15-OCT-2001 (9)

Type:

Lepomis macrochirus (Fish, fresh water) Species:

Exposure period: 96 hour(s)

Unit:

Analytical monitoring: yes

LC50: 11.5

Method: other: Standard Methods for the Examination of Water and

Wastewater, 13th ed., American Public Health Association

Year: GLP: no data

Test substance: other TS: Delac MOR Result: LC50 (96 hrs) = 11.5 ppm

The fish (1-3 inches in length) were obtained from a state Test condition:

licensed commercial fish hatchery and were retained for at least 10 days before use. The dilution water was natural pond water with a hardness expressed as 44 ppm CaCO3. A saturated aqueous extract (SAE) was made by allowing the material to sit in distilled water after vigorous shaking for at least 24 hours. The SAE was filtered and then tested. All tests were conducted at room temperature (22 degrees C), pH 6.5-8.5 and all DO's remained above 4mg/l at

all times during a test.

(1) valid without restriction Reliability:

Meets National standards method (AFNOR/DIN)

Critical study for SIDS endpoint Flag:

15-OCT-2001 (10)

- 12/60 -

Type: other: calculation

Species: other: Fish Exposure period: 96 hour(s)

Unit: mg/l Analytical monitoring: no

LC50: 1560.288

Method: other: ECOSAR v0.99e

Year: 1999 GLP: no

Test substance: other TS: molecular structure
Reliability: (2) valid with restrictions
Accepted calculation method

15-OCT-2001 (1)

Type: other: calculation Species: other: Saltwater Fish

Exposure period: 96 hour(s)

Unit: mg/l Analytical monitoring: no

LC50: 222.538

Method: other: ECOSAR v0.99e

Year: 1999 GLP: no

Test substance: other TS: molecular structure
Reliability: (2) valid with restrictions
Accepted calculation method

15-OCT-2001 (1)

Type: static

Species: Brachydanio rerio (Fish, fresh water)

Exposure period: 96 hour(s)

Unit: mg/l Analytical monitoring: no

LC0: 1 LC100: 5

Method:

Year: 1989 GLP: no

Test substance:

Remark: The substance was given in water and stirred for 24 h on a

magnetic stirrer: at all concentrations undissolved

particles remained in the medium.

24-APR-2001 (3)

Type: static

Species: Lepomis macrochirus (Fish, fresh water)

Exposure period: 96 hour(s)

Unit: mg/l Analytical monitoring: no

LC50: = 4.4

Method: other: Bionomics Lab protocol; see test conditions
Year: GLP: no data

Test substance: other TS: as prescribed by chapter 1 Monsanto

Remark: C.I. for 96 h LC50 = 3.6 - 5.6; 24 h LC50 > 7.5 mg/l; 48 h

LC50 = 6.0 mg/1

Test condition: carrier-acetone; 15l water; 10 fish/treatment; no aeration;

22 degrees C

24-APR-2001 (11)

- 13/60 -

Type: static

Species: Oncorhynchus kisutch (Fish, fresh water, marine)

Exposure period: 24 hour(s)

Unit: mg/l Analytical monitoring:

LC0: >= 10

Method:

Year: GLP: no

Test substance:

Remark: length: 5 - 10 cm; loss of equilibrium occured in 7 - 11 h;

only concentration tested

24-APR-2001 (12)

Type: static

Species: Oncorhynchus mykiss (Fish, fresh water)

Exposure period: 96 hour(s)

Unit: mg/l Analytical monitoring: no

NOEC: = .56 LC50: = 1.4

Method: OECD Guide-line 203 "Fish, Acute Toxicity Test"
Year: 1984 GLP: yes

Test substance: other TS: as prescribed by chapter 1 Monsanto

Remark: C.I. for LC50 = 1-1.8 mg/l; 24 h LC50 = 1.5 mg/l; 48 h LC50

= 1.4 mg/l

Test condition: hardness 40-45 mg/l CaCO3

24-APR-2001 (13)

Type: static

Species: Oncorhynchus mykiss (Fish, fresh water)

Exposure period: 96 hour(s)

Unit: mq/l Analytical monitoring: no

LC50: = 1.3

Method: other: Bionomics Lab protocol; see test conditions
Year: GLP: no data

Test substance: other TS: as prescribed by chapter 1 Monsanto

Remark: C.I. = 97-1.8 mg/l; 24 h LC50 = 5.3 mg/l; 48 h LC50 = 1.4

mg/l

Test condition: Carrier-acetone; 15 l water; 10 fish/treatment; length = 2.7

cm; no aeration; 12 degrees C

24-APR-2001 (11)

Type: static

Species: Oncorhynchus tschawytscha (Fish, fresh water, marine)

Exposure period: 24 hour(s)

Unit: mg/l Analytical monitoring:

LC0: >= 10

Method:

Year: GLP: no

Test substance:

Remark: length: 5 - 10 cm; only concentration tested

24-APR-2001 (12)

- 14/60 -

Type: static

Species: Ptychocheilus oregonensis (Fish, fresh water)

Exposure period: 24 hour(s)

Unit: mg/l Analytical monitoring:

LC0: >= 10

Method:

Year: GLP: no

Test substance:

Remark: length: 5 - 10 cm; only concentration tested

24-APR-2001 (12)

Type:

Species: Brachydanio rerio (Fish, fresh water)

Exposure period: 96 hour(s)

Unit: Analytical monitoring: no

Method:

Year: 1988 GLP: no

Test substance:

Remark: The substance was eluted in the concentration 1 g/l for 2 h,

filtered and the filtrate tested (DOC of the filtrate: 11

mg/1)

LCO at dilution 1:8 LC100 at dilution 1:2

24-APR-2001 (3)

4.2 Acute Toxicity to Aquatic Invertebrates

Type:

Species: Daphnia magna (Crustacea)

Exposure period: 48 hour(s)

Unit: mg/l Analytical monitoring: no

NOEC: = 1 EC50: = 4

Method: OECD Guide-line 202, part 1 "Daphnia sp., Acute

Immobilisation Test"

Year: 1984 GLP: no data

Test substance: other TS: Santocure MOR

Remark: C.I. for EC50 = 2.9-5.4; 24 h EC50 = 6.8 mg/l

Source: MonsantoBayer AG Leverkusen
Reliability: (1) valid without restriction

Guideline study

Flag: Critical study for SIDS endpoint

15-OCT-2001 (14)

- 15/60 -

Date: 15-OCT-2001 ID: 102-77-2 4. Ecotoxicity

Type:

Species: Daphnia magna (Crustacea)

Exposure period: 48 hour(s)

Unit: mq/1Analytical monitoring: no

NOEC: = 1 = 4.5 EC50:

other: EPA. Methods for Acute Toxicity Tests with Fish, Method:

Macroinvertebrates and Amphibians, EPA-660/3-75-009

Year: 1975 GLP: no data

Test substance: other TS: Santocure MOR

Remark: C.I. for 48 h EC50 = 3-6.8 mg/l; 24 h EC50 = 13 mg/l

Source: MonsantoBayer AG Leverkusen

Test condition: static, 250 ml water; 19 degrees C; 16 h light

Reliability: (1) valid without restriction

Guideline study

Critical study for SIDS endpoint Flaq:

15-OCT-2001 (15)

Type:

Species: other: Paratanytareum parthenogenetica

Exposure period: 48 hour(s)

Unit: mq/1Analytical monitoring: no

= 1.8 NOEC: = 5.3 EC50:

Method: other: EPA. Methods for Acute Toxicity Tests with Fish,

Macroinvertebrates and Amphibians, EPA-660/3-75-009

Year: 1975 GLP: yes

Test substance: other TS: Santocure MOR

Remark: C.I. for 48 h EC50 = 4.5 - 6.2 mg/l; 25 h EC50 = 8.6 mg/l

Source: MonsantoBayer AG Leverkusen

Test condition: static; 20 degrees C; 200 ml water; 16 h light

(1) valid without restriction Reliability:

GLP quideline study

Flag: Critical study for SIDS endpoint

15-OCT-2001 (16)

Type: other: calculation Species: Daphnia sp. (Crustacea)

Exposure period: 48 hour(s)

Unit: mq/1Analytical monitoring: no

LC50 : 1562.446

other: ECOSAR v0.99e Method:

Year: 1999 GLP: no

other TS: molecular structure Test substance: (2) valid with restrictions Reliability: Accepted calculation method

15-OCT-2001 (1)

- 16/60 -

Type: other: calculation

Species: Mysidopsis bahia (Crustacea)

Exposure period: 96 hour(s)

Unit: mq/1Analytical monitoring:

905.743 LC50 :

other: ECOSAR v0.99e Method:

Year: 1999 GLP: no

other TS: molecular structure Test substance: (2) valid with restrictions Reliability: Accepted calculation method

15-OCT-2001 (1)

4.3 Toxicity to Aquatic Plants e.g. Algae

Species: other algae: green algae

growth rate Endpoint: Exposure period: 96 hour(s)

Unit: mq/1Analytical monitoring: no

EC50: 923.219 ChV: 52.399

Method: other: ECOSAR v0.99e

1999 GLP: no Year:

Test substance: other TS: molecular structure (2) valid with restrictions Reliability: Accepted calculation method

Flag: Critical study for SIDS endpoint

15-OCT-2001 (1)

Species: other aquatic plant Endpoint: other: cell number

Exposure period: 96 hour(s)

Unit: mq/1Analytical monitoring:

EC50: 2

Method:

Year: GLP:

Test substance: 08-OCT-2001

other aquatic plant Endpoint: Species: other: chlorophyll a

Exposure period: 96

Unit: mq/1Analytical monitoring:

EC50: 2

Method:

Year: GLP:

Test substance: 08-OCT-2001

- 17/60 -

4.4 Toxicity to Microorganisms e.g. Bacteria

Type: aquatic

Species: activated sludge

Exposure period: 3 hour(s)

Unit: Analytical monitoring: no mg/l

EC50: > 10000

Method: ISO 8192 "Test for inhibition of oxygen consumption by

activated sludge"

Year: 1988 GLP: no

Test substance:

Remark: direct weight

(1) valid without restriction Reliability: Meets National standards method

15-OCT-2001 (3)

4.5 Chronic Toxicity to Aquatic Organisms

4.5.1 Chronic Toxicity to Fish

4.5.2 Chronic Toxicity to Aquatic Invertebrates

TERRESTRIAL ORGANISMS

4.6.1 Toxicity to Soil Dwelling Organisms

Type: other

Species: Eisenia fetida (Worm (Annelida), soil dwelling)

Endpoint:

Exposure period: 14 day other: mg/l Unit: LC50: 3110.605

Method: other:ECOSAR v0.99e

Year: 1999 GLP: no

Test substance: other TS: molecular structure Reliability: (2) valid with restrictions Accepted calculation method

15-OCT-2001 (1)

4.6.2 Toxicity to Terrestrial Plants

4.6.3 Toxicity to other Non-Mamm. Terrestrial Species

4.7 Biological Effects Monitoring

- 18/60 -

4.8 Biotransformation and Kinetics

-

4.9 Additional Remarks

-

- 19/60 -

5.1 Acute Toxicity

5.1.1 Acute Oral Toxicity

Type: LD50 Species: rat

Strain: Sprague-Dawley Sex: male/female

Number of

Animals: 16

Vehicle: other: corn oil Value: 12560 mg/kg bw

Method: other: Industrial BIO-TEST Laboratories, Inc.

Year: 1974 GLP: no

Test substance: other TS: OBTS, purity = 90-94%

Method: Industrial BIO-TEST Laboratories, Inc.

Initial scrrening was conducted to determine general level of

toxicity.

2 rats/sex/dose were administered the test material by gavage. The rats were then housed individually and observed for 14 days. Initial and final body weights, mortalities and

reactions were recorded. A necropsy examination was conducted

on all animals.

The acute oral median lethal dose was calculated using the techniques of Weil CS (1952); Thompson WR (1947) and Thompson

WR and Weil CS (1952).

Result: Adverse reactions observed at

4556 mg/kg: hyporeactivity and ruffed fur;

6834 mg/kg: as above plus salivation and labored breathing; 10250 mg/kg: as above plus muscular weakness, prostration,

diuresis

Dose level (mg/kg)	mortality rate	% mortality
4556 6834 10250 15380	0/4	0
	0/4	0
	1/4	25
	3/4	75

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Flag: Critical study for SIDS endpoint

15-OCT-2001 (17)

- 20/60 -

Type: LD50
Species: rat
Strain: Wistar
Sex: male/female

Number of

Animals: 20

Vehicle: other: propylene glycol

Value: > 10000 mg/kg bw

Method:

Year: GLP:

Test substance: other TS: Delac MOR

Method: The test material was given as a 33% (w/v) suspension in

propylene glycol to groups of 10 males and 10 femaels in a single dose of 30 ml/kg bw (10g test material/kg bw). The rats received feed and water ad libitum during the 14 day observation period. The rats were observed for intoxication

and mortality. All animals were necropsied.

Result: None of the treated animals showed any reaction upon

treatment. No deaths occurred during the observation period. Macroscopic examination did not reveal treatment-related

alterations.

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Flag: Critical study for SIDS endpoint

15-OCT-2001 (18)

Type: LD50 Species: rat

Strain:
Sex:
Number of
 Animals:
Vehicle:

Value: = 1980 mg/kg bw

Method:

Year: GLP:

Test substance:

Source: Bayer AG Leverkusen

06-APR-1992 (19)

Type: LD50 Species: rat

Strain:
Sex:
Number of
Animals:
Vehicle:

Value: > 7940 mg/kg bw

Method:

Year: GLP:

Test substance:

Remark: mortality: 0/7

Reliability: (2) valid with restrictions

15-OCT-2001 (20)

- 21/60 -

Type: LD50 Species: rat

Strain:
Sex:
Number of
 Animals:
Vehicle:

Value: > 5000 mg/kg bw

Method:

Year: GLP:

Test substance:

Remark: mortality: 0/10 Source: Bayer AG Leverkusen

06-APR-1992 (21)

Type: LD50 Species: mouse

Strain:
Sex:
Number of
 Animals:
Vehicle:

Value: = 1870 mg/kg bw

Method:

Year: GLP:

Test substance:

24-APR-2001 (22)

Type: LD50 Species: mouse

Strain:
Sex:
Number of
Animals:
Vehicle:

Value: = 1980 mg/kg bw

Method:

Year: GLP:

Test substance:

24-APR-2001 (23)

Type: LD50 Species: mouse

Strain:
Sex:
Number of
Animals:
Vehicle:

Value: = 4000 mg/kg bw

Method:

Year: GLP:

Test substance:

24-APR-2001 (24)

- 22/60 -

Type: LD100 Species: mouse

Strain:
Sex:
Number of
 Animals:
Vehicle:

Value: > 4000 mg/kg bw

Method:

Year: GLP:

Test substance:

24-APR-2001 (23)

Type: LDLo Species: mouse

Strain:
Sex:
Number of
Animals:
Vehicle:

Value: = 250 mg/kg bw

Method:

Year: GLP:

Test substance:

24-APR-2001 (23)

Type: other: LD Species: rabbit

Strain:
Sex:
Number of
 Animals:
Vehicle:

Value: > 3980 mg/kg bw

Method:

Year: GLP:

Test substance:

24-APR-2001 (20)

5.1.2 Acute Inhalation Toxicity

Type: LC50 Species: rat

Strain:

Sex: male

Number of

Animals: 10

Vehicle:

Exposure time: 1 hour(s)
Value: > 151 mg/l

Method: other: according to the technique specified in the Regulations

(Federal Register, August 12, 1961)

Year: 1961 GLP: no

Test substance: other TS: OBTS; purity not noted

- 23/60 -

Result: Two rats died on the 4th or 10th day post-exposure. No other

mortalities occurred.

During the one-hour exposure, animals generally exhibited excessive preening and masticatory movements, excessive salivation, and occasional periods of excited activity.

Signs of toxicity prior to the mortalities were depression, depresed righting and placement reflexes, ataxia (1 animal-4th day post exposure) and diarrhea stains of 2 day duration (1 animal -10th day post exposure).

(1 allillial -10th day post exposure).

Surviving rats exhibited normal appearance and behavior

throughout the 14-day observation period.

Test condition: Sample was ground in a blender and filtered through a

40-mesh screen; then generated as a dust for exposure.

Reliability: (1) valid without restriction

Meets National standards method

Flag: Critical study for SIDS endpoint

15-OCT-2001 (25)

Type: LC0 Species: rat

Strain:

Sex: male/female

Number of

Animals: 20

Vehicle: other: undiluted powder

Exposure time: 4 hour(s)
Value: ca. .09 mg/l

Method: other: Industrial Bio-Test Laboratories protocol

Year: 1974 GLP: no

Test substance: as prescribed by 1.1 - 1.4

Method: Ten albino rats were exposed to the dust of the test

material for four hours in a 70 liter chamber. The

concentration was determined by sampling the test atmosphere in the breathing zone of the animals, collected on a glass fiber filter. The average filter concentration was 0.09 mg/l air. The exposed animals and 10 untreated controls were

observed for 14 days.

Result: There were no deaths during the exposure or 14-day observation

period. All rats in the treated and control groups gained weight during the study. There were no gross pathologic findings attributable to inhalation of the test material. Gross pathologic changes in test and untreated control

animals were essentially the same.

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Flag: Critical study for SIDS endpoint

15-OCT-2001 (26)

- 24/60 -

5.1.3 Acute Dermal Toxicity

Type: LD50 Species: rabbit

Strain: New Zealand white

Sex: male/female

Number of

Animals:

Vehicle: other: neat Value: > 3000 mg/kg bw

Method:

Year: GLP: no

Test substance: other TS: OBTS

Method: Young adult New Zealand albino rabbits were acclimated and

> examined prior to testing. Twenty-four hours prior to testing, the backs of the animals (approx 30% total body surface) were shaved free of hair. The material was applied at the highest reasonable dosage and the test site was covered with plastic sheeting. After 24 hours the plastic sheeting and residual material were removed. The animals

were examined for local skin reactions, behavioral

abnormalities and mortality for 14 days. Initial, 7 and 14 day body weights were recorded. A necropsy was conducted on

all animals.

Result: mortality = 0/4

Test condition: The test material was applied as received to abraded,

pre-moistened skin.

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Critical study for SIDS endpoint Flaq:

15-OCT-2001 (17)

LD50 Type: Species: rabbit

Sex: Number of

Strain:

Animals: Vehicle:

Value: > 7940 mg/kg bw

Method:

Year: GLP:

Test substance:

24-APR-2001 (27)

- 25/60 -

Type: other: LD Species: rabbit

Strain:
Sex:
Number of
 Animals:
Vehicle:

Value: > 3980 mg/kg bw

Method:

Year: GLP:

Test substance:

24-APR-2001 (28)

5.1.4 Acute Toxicity, other Routes

Type: LD50 Species: mouse

Strain:
Sex:
Number of
Animals:
Vehicle:

Route of admin.: i.p.

Value: ca. 100 - 200 mg/kg bw

Method:

Year: GLP:

Test substance:

24-APR-2001 (29)

Type: LD50 Species: mouse

Strain:
Sex:
Number of
 Animals:
Vehicle:

Route of admin.: i.p.

Value: = 100 mg/kg bw

Method:

Year: GLP:

Test substance:

24-APR-2001 (30)

- 26/60 -

Type: LD50 Species: mouse

Strain:
Sex:
Number of
 Animals:
Vehicle:

Route of admin.: other: no data Value: = 3150 mg/kg bw

Method:

Year: GLP:

Test substance:

Remark: sex: male

24-APR-2001 (31)

Type: LD50 Species: mouse

Strain:
Sex:
Number of
 Animals:
Vehicle:

Route of admin.: other: no data Value: = 3000 mg/kg bw

Method:

Year: GLP:

Test substance:

Remark: sex: female

24-APR-2001 (31)

5.2 Corrosiveness and Irritation

5.2.1 Skin Irritation

Species: rabbit

Concentration: undiluted

Exposure: Occlusive

Exposure Time: 24 hour(s)

Number of Animals:

PDII: .8

Result: slightly irritating
EC classificat.: not irritating
Method: Draize Test

Year: GLP: no data

Test substance: other TS: OBTS; purity not noted

Method: dose: 0.5 g undiluted applied to intact and abrdaded skin;

occluded for 24 hours.

Result: sample tested Primary Irritation Score

OBTS 1-A 0.1
OBTS 1-B 0.2
OBTS 1-C 0.1
OBTS 1-D 0.8

- 27/60 -

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Flag: Critical study for SIDS endpoint

15-OCT-2001 (17)

Species: rabbit

Concentration:

Exposure: Exposure Time: Number of

Animals: 12

PDII:

Result: slightly irritating

EC classificat.:

Method:

Year: GLP:

Test substance: other TS: Delac MOR

The test substance caused very slight skin irritation in 2 of Result:

12 rabbits.

15-OCT-2001 (32)

Species: rabbit

Concentration:

Exposure: Exposure Time: Number of Animals: PDII:

Result:

not irritating

EC classificat.:

Method: other: 24 hours

Year: GLP:

Test substance:

Remark: score: 0.6/8.0, practically non-irritating

15-OCT-2001 (27)

rabbit Species:

Concentration:

Exposure: Exposure Time: Number of Animals: PDII:

Result: not irritating

EC classificat.:

Method: other: 500 mg/animal, 2 animals, onto the skin of the ear for

24 h (semi-occlusive), test substance removed with water at

the end of exposure; post exposure period of 7 d

Year: GLP:

Test substance:

Remark: the same procedure with a test substance of different

- 28/60 -

origin came to the result: slight skin irritant

24-APR-2001 (33)

Species: rabbit

Concentration:

Exposure:
Exposure Time:
Number of
Animals:
PDII:

Result: not irritating

EC classificat.:

Method: other: test substance was applied to the clipped, intact skin

(occlusive) and removed after 24 h with water, observation over several days, the data were scored according to the

method of Draize

Year: GLP:

Test substance:

24-APR-2001 (28)

Species: guinea pig

Concentration:

Exposure:
Exposure Time:
Number of
Animals:
PDII:

Result:

EC classificat.:

Method: other: 0.5, 2, 5 and 10 % being tested to determine the

threshold irritation concentration, application over 24 h (semi-oclusive), readings were performed at 1, 24 and 48 h $\,$

after removal

Year: GLP:

Test substance:

Remark: result: 5 and 10 % produced irritant reactions

Source: Bayer AG Leverkusen

15-OCT-1993 (34)

- 29/60 -

5.2.2 Eye Irritation

Species: rabbit

Concentration: undiluted Dose: 100 other: mg

Exposure Time:

Comment: not rinsed

Number of

Animals: 24

Result: moderately irritating

EC classificat.: irritating
Method: Draize Test

Year: GLP: no data

Test substance: other TS: OBTS; purity not noted Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Flag: Critical study for SIDS endpoint

15-OCT-2001 (17)

Species: rabbit

Concentration:

Dose:

Exposure Time: Comment:

Number of

Animals: 6

Result: moderately irritating

EC classificat.: irritating

Method:

Year: GLP:

Test substance: other TS: Delac MOR

Result: The test substance caused moderate ocular lesions in 4 of 6

rabbits.

15-OCT-2001 (32)

Species: rabbit

Concentration:

Dose:

Exposure Time:
Comment:
Number of
Animals:

Result: slightly irritating

EC classificat.:

Method: other: 24 hours

Year: GLP:

Test substance:

Remark: score: 6.0/110.0

15-OCT-2001 (27)

- 30/60 -

Species: rabbit

Concentration:

Dose:

Result:

Exposure Time:
Comment:
Number of
Animals:

EC classificat.:

Method: other: 100mg/24h

Year: GLP:

Test substance:

Remark: result: moderate

24-APR-2001 (35)

Species: rabbit

Concentration:

Dose:

Exposure Time:
Comment:
Number of

Animals:

Result: slightly irritating

EC classificat.:

Method: other: 50 mg/animal, 2 animals, instillation into the

conjunctival sac, post exposure period of 7 d

Year: GLP:

Test substance:

Remark: the same procedure with a test substance of different

origin came to the result: moderate eye irritant

24-APR-2001 (33)

Species: rabbit

Concentration:

Dose:

Exposure Time:
Comment:
Number of

Number of Animals:

Result: slightly irritating

EC classificat.:

Method: other: finely ground test substance were placed in the

conjunctival sac, the eyes were rinsed after 24 h, the data

scored according to the method of Draize

Year: GLP:

Test substance:

24-APR-2001 (28)

- 31/60 -

5.3 Sensitization

Type: other: see remarks

Species: guinea pig

Number of
 Animals:
Vehicle:

Result: sensitizing

Classification:

Method: other: a modification of Buehler's method

Year: GLP:

Test substance:

Remark: cross-sensitivity: 8/10 sensitized animals showed a

positive reaction with MBT

15-OCT-2001 (34)

Type: other: see remarks

Species: guinea pig

Number of Animals: Vehicle:

Result: sensitizing

Classification:

Method: other: closed epicutaneous test (no further information)

Year: GLP:

Test substance:

Remark: result: 8/12 animals positive (challenge with 0.5M)
1/12 animals positive (challenge with 0.05M)

15-OCT-2001 (36)

Type: Patch-Test Species: human

Number of
 Animals:
Vehicle:
Result:

Classification:

Method:

Year: GLP:

Test substance:

Remark: 4/31 rubber contact dermatitis patients had a positive patch

test result

15-OCT-2001 (36)

- 32/60 -

Type: Patch-Test Species: human

Number of
Animals:
Vehicle:
Result:

Classification:

Method:

Year: GLP:

Test substance:

Remark: 17/17 subjects allergic to MBT were positive in the test with MBS; the test with MBS was negative in 20 controls

15-OCT-2001 (37)

Type: Patch-Test Species: human

Number of
Animals:
Vehicle:
Result:

Classification:

Method:

Year: GLP:

Test substance:

Remark: 24/49 individuals showed a positive patch test reaction during the induction phase (for the challenge applications,

petrolatum alone was applied in lieu of test sample)

24-APR-2001 (38)

Type: Patch-Test Species: human

Number of
Animals:
Vehicle:
Result:

Classification:

Method:

Year: GLP:

Test substance:

Remark: Repeat patch testing with 1 or 10 % Santocure MOR resulted

in sensitization reactions in 7/24 and 28/31 subjects, respectively. Recrystallized Santocure MOR showed only 1/20

individuals being affected.

15-OCT-2001 (39)

- 33/60 -

Type: Patch-Test

Species: human

Number of Animals: Vehicle: Result:

Classification:

Method:

Year: GLP:

Test substance:

Remark: Evaluation of Santocure MOR, mercaptobenzothiazole,

morpholine and two oxidized impurities, the sulfonamide and the sulfenamide, were tested in vitro with lymphocytes from

sensitized individuals to determine if DNA

synthesis/stimulation would result. The results showed the oxidized contaminants to be primarily responsible for the

sensitization response.

15-OCT-2001 (40)

Type: other: in vitro lymphocyte assay

Species:
Number of
 Animals:
Vehicle:
Result:

Classification:

Method:

Year: GLP:

Test substance:

Remark: An in vitro lymphocyte assay was performed to determine if

Santocure MOR or process contaminants might possess the ability to induce delayed type hypersensitivity. Based on the data, it appears that Santocure MOR is not a sensitizer,

but two potential process contaminants may be.

24-APR-2001 (41)

- 34/60 -

5.4 Repeated Dose Toxicity

Species: rat Sex:

Strain:

Route of admin.: oral feed

Exposure period: 4 w

Frequency of
 treatment:
Post. obs.
 period:

Doses: 100, 200, 500, or 1000 mg/kg bw/d

Control Group:

NOAEL: 200 mg/kg bw LOAEL: 500 mg/kg bw

Method:

Year: GLP:

Test substance: other TS: Santocure MOR

Result: Body weight reductions were noted in animals at 500 and 1000

mg/kg. Increased liver and kidney weights were observed in

the high-dose males

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Flag: Critical study for SIDS endpoint

15-OCT-2001 (42)

Species: rat Sex: male/female

Strain: other: Charles River CD strain

Route of admin.: oral feed Exposure period: 113 weeks

Frequency of

treatment: daily

Post. obs. period:

Doses: 5, 50, 400 mg/kg bw/d

Control Group: yes

NOAEL: 5 mg/kg bw LOAEL: 50 mg/kg bw

Method: other: according to Hazelton Laboratories Europe Ltd.

Year: 1982 GLP: no data

Test substance: other TS: Santocure MOR

Result: 50 and 400 mg/kg bw: reductions in body weight gain and

food consumption; dose related increases in kidney and liver

weights; no other evidence of chronic toxicity

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Flag: Critical study for SIDS endpoint

15-OCT-2001 (43)

- 35/60 -

Species: rat Sex:

Strain:

Route of admin.: inhalation

Exposure period: 4 w

Frequency of

treatment: 6 h/d at 5 d/w

Post. obs. period:

Doses: 4.4, 9.8, and 10.2 mg/m3

Control Group:

NOAEL: 9.8 mg/m³

Method:

Year: GLP:

Test substance:

Result: Santocure MOR caused slight irratation in exposed animals.

Slight body weight reductions and reductions in lung weights

were observed in males of high-exposure group.

Histopathological examination revealed no alterations in the

tissues of the high-exposure group animals. Slight

depressions in blood glucose and elevations in SGOT values were found in animals at all exposure levels, but these findings were not associated with the presence of any tissue

lesions.

The no effect level for this study was considered to be 9.8

mg/m3.

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Flag: Critical study for SIDS endpoint

15-OCT-2001 (44)

Species: rabbit Sex:

Strain:

Route of admin.: dermal Exposure period: 21 d

Frequency of
 treatment:
Post. obs.
 period:

Doses: 125, 500, 2000

Control Group:

NOAEL: 2000

Method:

Year: GLP:

Test substance:

Result: Repeated applications of Santocure MOR produced no evidence

of toxicity related to test material administration. Only a

slight degree of dermal irritation was noted.

15-OCT-2001 (44)

- 36/60 -

Species: mouse Sex: male/female

Strain: other: Slc:ddY
Route of admin.: oral feed
Exposure period: 3 months

Frequency of

treatment: no data

Post. obs.

period: no data

Doses: 0.012, 0.046, 0.188, 0.75 % (18, 69, 282, 1125 mg/kg bw)

Control Group: no data specified NOAEL: 282 mg/kg bw LOAEL: 1125 mg/kg bw

Method:

Year: GLP:

Test substance:

Result: 0.75 % group: decreased body weight gain, slightly

increased levels of GPT and kidney weights (no further

information)

15-OCT-2001 (31)

Species: rat Sex: no data

Strain: no data
Route of admin.: inhalation

Exposure period: 15 d

Frequency of

treatment: 2h/d

Post. obs. period:

Doses: 0.3 - 0.4 mg/l Control Group: no data specified

Method:

Year: GLP:

Test substance:

Remark: no detailed information; evaluation impossible

Result: no change in body weight; the working of the nervous system

was affected

Reliability: (4) not assignable

Documentation insufficient for assessment

15-OCT-2001 (23)

- 37/60 -

Species: rat Sex: male

Strain: Sprague-Dawley

Route of admin.: gavage Exposure period: 56 d

Frequency of

treatment: daily, 7d/w

Post. obs.

period: 2w

Doses: 125, 250, 500 mg/kg bw Control Group: yes, concurrent vehicle

Method:

Year: GLP:

Test substance:

Result: mean body weight and mean body weight gain did not reveal

any significant effect in any of the dose groups;

tissue-to-body weight ratios were normal except an increase in the stomach/body weight ratio (125, 250 mg/kg bw); no abnormalities in selected animals during gross pathological

examination

24-APR-2001 (45)

Species: mouse Sex: male/female

Strain: other: Slc:ddY
Route of admin.: oral feed
Exposure period: 21 months

Frequency of

treatment: no data

Post. obs.

period: no data

Doses: 0.01, 0.1, 1 %
Control Group: no data specified
NOAEL: 150 mg/kg bw
LOAEL: 1500 mg/kg bw

Method:

Year: GLP:

Test substance:

Remark: doses: 15, 150, 1500 mg/kg bw remarks: see also chapter 5.7

Result: 1 % group: decreased body weight gain (no further

information)

15-OCT-2001 (31)

- 38/60 -

Date: 15-OCT-2001
5. Toxicity

Date: 15-OCT-2001

Species: mouse Sex: no data

Strain: other: Slc:ddY

Route of admin.: dermal Exposure period: 21 months

Frequency of

treatment: no data

Post. obs.

period: no data

Doses: 10 % suspended in olive oil

Control Group: no data specified

NOAEL: 10 %

Method:

Year: GLP:

Test substance:

Result: no compound related adverse effects (no further information) 15-OCT-2001 (31)

Species: rabbit Sex: no data

Strain: no data

Route of admin.: oral unspecified

Exposure period: 3.5 months

Frequency of

treatment: on alternative days (2 months), daily (1.5 months)

Post. obs.

period: no data
Doses: 20 mg/kg bw
Control Group: no data specified

Method:

Year: GLP:

Test substance:

Remark: no detailed information; evaluation impossible

Result: general conditions were unchanged; the pigmentary function (?) of the liver was affected; changes in the liver, kidneys

(:) Of the liver was affected, changes in the liver, kidneys

and lung were observed

24-APR-2001 (23)

5.5 Genetic Toxicity 'in Vitro'

Type: Ames test

System of

testing: Salmonella typhimurium TA 98, TA 100, TA 1535, TA 1537, TA

1538

Concentration: <= 500 ug/ml
Cytotoxic Conc.: >= 500 ug/plate

Metabolic

activation: with and without

Result: negative

Method: OECD Guide-line 471 "Genetic Toxicology: Salmonella

thyphimurium Reverse Mutation Assay"

Year: GLP: yes

Test substance: other TS: Three sample comparison (two comercial: OBTS and

NOBS; and OBTS 4X recrystallized sample)

Reliability: (1) valid without restriction

GLP guideline study

- 39/60 -

Critical study for SIDS endpoint

15-OCT-2001 (46)

Type: Mammalian cell gene mutation assay

System of

testing: CHO cells

Concentration: 0.1, 0.3, 1.0, 3.0, 10, 30, 50, 100, 150, 300 ug/ml

Cytotoxic Conc.: >=150 ug/ml (with and without activation)

Metabolic

with and without activation:

Result: negative

Method: OECD Guide-line 476 "Genetic Toxicology: In vitro Mammalian

Cell Gene Mutation Tests"

Year: GLP: yes Test substance: other TS: Commercial and purified OBTS

Reliability: (1) valid without restriction

GLP guideline study

Flaq: Critical study for SIDS endpoint

15-OCT-2001 (47)

Type: Mammalian cell gene mutation assay

System of

testing: mouse lymphona cells L 5158 TK+/- Concentration: <=50.0 mg/plate

Cytotoxic Conc.: > 50 ug/ml

Metabolic

activation: with and without

Result: positive

Method: OECD Guide-line 476 "Genetic Toxicology: In vitro Mammalian

Cell Gene Mutation Tests"

Year: 1981 GLP: yes

other TS: Three sample comparison (two comercial: OBTS and Test substance:

NOBS; and OBTS 4X recrystallized sample)

An analysis of small vs. large colonies was not made, however Result:

> because the lower-limit cutoff on the colony counter is 0.3mm, it is assumed that the mutation response was mainly due to

large colonies.

The assay was positive for all three test substances under

conditions of metabolic activation.

(1) valid without restriction Reliability:

GLP guideline study

Flaq: Critical study for SIDS endpoint

15-OCT-2001 (46)

- 40/60 -

Date: 15-OCT-2001
5. Toxicity

Date: 15-OCT-2001

Type: Sister chromatid exchange assay

System of

testing: CHO cells

Concentration: 1,5,10,20,40 ug/ml (non-activated) 5,10,15,30,60 ug/ml

(activated)

Cytotoxic Conc.: >= 50 ug/ml

Metabolic

activation: with and without

Result: negative

Method: OECD Guide-line 479 "Genetic Toxicology: In vitro Sister

Chromatid Exchange Assay in Mammalian Cells"

Year: 1984 GLP: yes Test substance: other TS: OBTS commercial and purified

Reliability: (1) valid without restriction

GLP guideline study

Flag: Critical study for SIDS endpoint

15-OCT-2001 (48)

Type: Cytogenetic assay

System of

testing: CHO cells
Concentration: <= 10 mg/ml</pre>

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative Method: other

Year: 1979 GLP:

Test substance: other TS: purity = 90-95 %

Remark: GLP: Signed Quality Assurance Inspection Statement

Reliability: (1) valid without restriction

Meets generally accepted scientific method and is described in

sufficient detail

Flag: Critical study for SIDS endpoint

15-OCT-2001 (49)

Type: other: Cell Transformation Assay

System of

testing: BALB / 3T3 Cell line

Concentration: <= 30.0 mg/l

Cytotoxic Conc.:

Metabolic

activation: without
Result: negative
Method: other

Year: 1981 GLP:

Test substance: other TS: Three sample comparison (two comercial: OBTS and

NOBS; and OBTS 4X recrystallized sample)

Remark: GLP: Signed Quality Assurance Inspection Statement

Reliability: (1) valid without restriction

Meets generally accepted scientific method and is described in

sufficient detail

Flag: Critical study for SIDS endpoint

15-OCT-2001 (50)

- 41/60 -

Type: other: DNA repair assay

System of

testing: E. coli W3110 (pol A+) E. coli W3078 (pol A-)

Concentration: <= 1.0 mg/plate</pre>

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative Method: other

Year: 1979 GLP:

Test substance: other TS: 90-95 %

Remark: GLP: Signed Quality Assurance Inspection Statement

Reliability: (1) valid without restriction

Meets generally accepted scientific method and is described in

sufficient detail

Flag: Critical study for SIDS endpoint

15-OCT-2001 (51)

Type: Ames test

System of

testing: Salmonella typhimurium TA 98, TA 100, TA 1535, TA 1537, TA

1538

Concentration: <= 5.0 mg/plate</pre>

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method: OECD Guide-line 471 "Genetic Toxicology: Salmonella

thyphimurium Reverse Mutation Assay"

Year: 1979 GLP: Test substance: other TS: OBTS; purity =90-95 %

Remark: GLP: Signed Quality Assurance Inspection Statement

Reliability: (1) valid without restriction

Meets generally accepted scientific method and is described in

sufficient detail

15-OCT-2001 (49)

Type: Ames test

System of

testing: Salmonella typhimurium TA 98, TA 100, TA 1535, TA 1537, TA

1538

Concentration: <= 1000 ug/plate
Cytotoxic Conc.: >= 100 ug/plate

Metabolic

activation: with and without

Result: negative

Method: OECD Guide-line 471 "Genetic Toxicology: Salmonella

thyphimurium Reverse Mutation Assay"

Year: 1982 GLP: no data

Test substance: other TS: OBTS; purity > 99 %

Method: The standard Ames protocol was followed with the following

exception: instead of duplicate plates for the positive controls, in trial #1, one of the plates contained twice the

standard dose.

Reliability: (1) valid without restriction

- 42/60 -

Meets generally accepted scientific method and is described in

sufficient detail

15-OCT-2001 (52)

Type: Bacterial gene mutation assay

System of

testing: Salmonella typhimurium TA 100, TA 98

Concentration: Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method:

Year: GLP:

Test substance:

15-OCT-2001 (53)

Type: Ames test

System of

testing: Salmonella typhimurium TA 98, TA 100

Concentration: Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method:

Year: GLP:

Test substance:

15-OCT-2001 (54)

Type: Ames test

System of

testing: Salmonella typhimurium TA 1535, TA 1537, TA 98, TA 100

Concentration:
Cytotoxic Conc.:

Metabolic

activation: with Result: negative

Method:

Year: GLP:

Test substance:

15-OCT-2001 (55) (56)

- 43/60 -

Date: 15-OCT-2001
5. Toxicity

Date: 15-OCT-2001

Type: Ames test

System of

testing: Salmonella typhimurium TA 98, TA 100, TA 1535, TA 1537, TA

1538

Concentration:
Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative Method: other

Year: 1976 GLP:

Test substance:

15-OCT-2001 (57)

Type: Mammalian cell gene mutation assay

System of

testing: CHO cells
Concentration: 2.5 - 50 ug/ml
Cytotoxic Conc.: >=40 ug/ml

Metabolic

activation: with and without

Result: negative

Method: OECD Guide-line 476 "Genetic Toxicology: In vitro Mammalian

Cell Gene Mutation Tests"

Year: 1986 GLP:

Test substance: other TS: OBTS commercial: purity = 94.12%, purified: purity

>= 99%

Remark: GLP: Signed Quality Assurance Inspection Statement

commercial and purified sample
(1) valid without restriction

Reliability: (1) valid without restric Guideline study

15-OCT-2001

Type: Mammalian cell gene mutation assay

System of

testing: mouse lymphoma cells L5178Y TK+/-

Concentration: <= 50.0 mg/plate</pre>

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: positive Method: other

Year: 1979 GLP:

Test substance: other TS: 90-95 % purity

Remark: GLP: Signed Quality Assurance Inspection Statement

result: significant increases in the mutation frequency were

(58)

observed in the absence and presence of rat liver S9

metabolic activation

Reliability: (1) valid without restriction

Meets generally accepted scientific method and is described in

sufficient detail

15-OCT-2001 (49)

- 44/60 -

Type: Mammalian cell gene mutation assay

System of

testing: mouse lymphoma cells L5178Y TK+/-

Concentration:
Cytotoxic Conc.:

Metabolic

activation: with and without

Result: positive

Method:

Year: GLP:

Test substance:

Remark: result: significant increases in the mutation frequeny were

only observed in the presence of rat liver S9 metabolic

activation

15-OCT-2001 (59)

Type: other: Cell transformation assay

System of

testing: BALB/3T3 Cell line

Concentration: <= 35.0 mg/l

Cytotoxic Conc.:

Metabolic

activation: without
Result: positive
Method: other

Year: 1979 GLP:

Test substance: other TS: 90-95 %

Remark: GLP: Signed Quality Assurance Inspection Statement

Reliability: (1) valid without restriction

Meets generally accepted scientific method and is described in

sufficient detail

15-OCT-2001 (49)

Type: Yeast gene mutation assay

System of

testing: Saccharomyces cerevisiae D4

Concentration: <= 1.0 mg/plate</pre>

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method:

Year: 1979 GLP:

Test substance: other TS: 90-95 %

Remark: GLP: Signed Quality Assurance Inspection Statement

15-OCT-2001 (51)

- 45/60 -

Type: Yeast gene mutation assay

System of

testing: Saccharomyces cerevisiae D4

Concentration:
Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method:

Year: GLP:

Test substance:

15-OCT-2001 (57)

Type: other: DNA repair assay

System of

testing: E. coli W1310 (polA+), E. coli p3478 (polA-)

Concentration: Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method: OECD Guide-line 472 "Genetic Toxicology: Escherichia coli

Reverse Mutation Assay"

Year: GLP:

Test substance: other TS: Three sample comparison (two comercial: OBTS and

NOBS; and OBTS 4X recrystallized sample)

Reliability: (1) valid without restriction

Guideline study

15-OCT-2001 (60)

Type: other: DNA repair suspension assay

System of

testing: E. coli W3110 (pol A+) E. coli P3078 (pol A-)

Concentration: <= 2.5 mg/plate</pre>

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: positive Method: other

Year: 1981 GLP:

Test substance: other TS: 90-95 % purity

Remark: GLP: Signed Quality Assurance Inspection statement

Reliability: (1) valid without restriction

Meets generally accepted scientific method and is described in

sufficient detail

15-OCT-2001 (49)

- 46/60 -

5.6 Genetic Toxicity 'in Vivo'

Type: Dominant lethal assay

Species: rat Sex: male

Strain: Sprague-Dawley

Route of admin.: gavage Exposure period: 56d

Doses: 125, 250, 500 mg/kg bw

Result: negative Method: other

Year: 1980 GLP: yes

Test substance: other TS: 90-95 % purity Remark: see also chapter 5.4

Result: no dominant lethal mutations
Reliability: (1) valid without restriction

GLP study; Meets generally accepted scientific method and is

described in sufficient detail Critical study for SIDS endpoint

15-OCT-2001 (45)

Type: Dominant lethal assay

Species: rat Sex: male/female

Strain:

Flag:

Route of admin.: oral unspecified

Exposure period: up to 3d

Doses: 200 mg/kg bw

Result: Method:

Year: GLP:

Test substance:

Remark: no further information

Result: An increase in total embryonic mortality in treated females

and in females mated with treated males.

15-OCT-2001 (61)

Type: unspecified

Species: Drosophila melanogaster Sex: no data

Strain:

Route of admin.: other
Exposure period: no data
Doses: no data

Result: Method:

Year: GLP:

Test substance:

Result: Not described in detail.

Opinion of the author: MBS showed a weak mutagenic activity.

15-OCT-2001 (62)

- 47/60 -

5.7 Carcinogenicity

Species: mouse Sex:

Strain:

Route of admin.: gavage Exposure period: 79 w

Frequency of treatment:
Post. obs.
period:

Doses: 90 mg/kg/d Result: negative

Control Group:

Method:

Year: GLP:

Test substance: other TS: N-oxydiethylene-2-benzothiazole sulfenamide (NOBS)

Result: No adverse effects were reported, and no statistically

significant increase in tumor incidences were observed in

the study.

25-APR-2001 (63)

Species: rat Sex: no data

Strain: no data
Route of admin.: oral feed
Exposure period: 2 years

Frequency of treatment:
Post. obs.
period:

Doses: 5, 50, 400 mg/kg bw/d

Result: negative

Control Group: no data specified

Method:

Year: GLP:

Test substance:

Remark: no further information; see also chapter 5.4

Result: no evidence of oncogenicity

15-OCT-2001 (64)

- 48/60 -

Species: mouse Sex: male/female

Strain: other: Slc:ddY
Route of admin.: oral feed
Exposure period: 21 months

Frequency of

treatment: no data

Post. obs.

Result: negative

Control Group: no data specified

Method:

Year: GLP:

Test substance:

Remark: doses: 15, 150, 1500 mg/kg bw remarks: see also chapter 5.4

Result: histopathologically no compound related nonneoplastic or

neoplastic lesions (no further information)

15-OCT-2001 (31)

Species: mouse Sex: male/female Strain: other: (C57BL/6 x C3H/Anf)Fl and (C57BL/6 x AKR)Fl

Route of admin.: oral unspecified

Exposure period: 18 months

Frequency of

treatment: daily

Post. obs.

period: no

Doses: 464 mg/kg bw (days 7-28 of age); 1492 ppm (after 28 days of

age)

Result: negative

Control Group: yes

Method:

Year: GLP:

Test substance:

Remark: doses: 1492 ppm = dosage in diet = approx. 224 mg/kg bw no significant indication of tumorigenicity after oral ad-

ministration

25-APR-2001 (65)

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Species: mouse Sex: male/female Strain: other: (C57BL/6 x C3H/Anf)Fl abd (C57BL/6 x AKR)Fl

Route of admin.: s.c. Exposure period: 1d

Frequency of

treatment: once

Post. obs.

period: 18 months
Doses: 464 mg/kg bw
Result: negative
Control Group: yes

Method:

Year: GLP:

Test substance:

Result: no significant indication of tumorigenicity

15-OCT-2001 (66)

Species: mouse Sex:

Strain:

Route of admin.: s.c.

Exposure period: Frequency of

treatment: single injection

Post. obs.

period: 78 w

Doses: 1000 mg/kg Result: negative

Control Group:

Method:

Year: GLP:

Test substance: other TS: N-oxydiethylene-2-benzothiazole sulfenamide (NOBS)

Result: No adverse effects were reported, and no statistically

significant increase in tumor incidences were observed in

the study.

15-OCT-2001 (63)

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5.8 Toxicity to Reproduction

Type:

Species: rat Sex: male

Strain: Sprague-Dawley

Route of admin.: gavage Exposure Period: 56d

Frequency of

treatment: daily, 7d/w

Duration of test:

Doses: 125, 250, 500 mg/kg bw Control Group: yes, concurrent vehicle

Method:

Year: GLP:

Test substance:

Remark: see also chapter 5.4

post observation period: 2w

Result: pregnancy rates of the dose groups were comparable to those

of the controls

25-APR-2001 (45)

Type:

Species: rat Sex: male/female

Strain:

Route of admin.: gavage Exposure Period: 3d

Frequency of

treatment: 1st and 3rd days of estrus (female); twice an interval of 3d

Duration of test:

Doses: 200 mg/kg bw

Control Group: yes

Method:

Year: GLP:

Test substance:

Remark: The object of the investigations was to study the level

of the embryonic mortality (EM).

post observation period: till 19th day of pregnancy no visible signs of poisoning; changes in the estrous

cycle; delay to conceptions; decreased fetus wieghts; increased total embryonic mortality (EM) with normal

postimplantation EM

25-APR-2001 (67)

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5.9 Developmental Toxicity/Teratogenicity

Species: rat Sex: female

Strain: other: Charles River Route of admin.: oral unspecified Exposure period: 6 - 15 gestation day

Frequency of

treatment: daily

Duration of test: until 20 gestation day Doses: 100, 300, 1000 mg/kg bw

Control Group: yes

NOAEL Maternalt.: 300 mg/kg bw NOAEL Teratogen.: 1000 mg/kg bw

Method:

Year: GLP:

Test substance: other TS: Santocure MOR

Remark: Maternal toxicity was noted at the highest dose level.

Result: no teratogenic response was observed (no further

information)

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Flag: Critical study for SIDS endpoint

15-OCT-2001 (68)

Species: rat Sex: female

Strain: no data Route of admin.: oral feed

Exposure period: day 0 through 20 of gestation

Frequency of
 treatment:
Duration of test:

Doses: 0.02, 0.5 % (13, 270 mg/kg bw/d)

Control Group: yes

NOAEL Maternalt.: 270 mg/kg bw NOAEL Teratogen.: 270 mg/kg bw

Method:

Year: GLP:

Test substance: other TS: Santocure MOR Remark: no further information

Result: no teratogenic, fetotoxic or maternally toxic effects

25-APR-2001 (27)

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Species: rat Sex: female

Strain:

Route of admin.: gavage Exposure period: 2d

Frequency of

treatment: 4th and 11th days of pregnancy
Duration of test: till 19th day of pregnancy

Doses: 200 mg/kg bw Control Group: other: yes

Method:

Year: GLP:

Test substance:

Remark: The object of the investigations was to study the level of

the embryonic mortality (EM).

Result: no visible signs of poisoning; decreased fetus weights; in-

creased total embryonic mortality (EM) with normal

postimplantation EM

25-APR-2001 (67)

Species: rat Sex: female

Strain: other: wistar Route of admin.: oral feed

Exposure period: day 0 of gestation to day 21 postparturition

Frequency of
 treatment:
Duration of test:

Doses: 0.02, 0.5 %

Control Group: yes

Method:

Year: GLP:

Test substance:

Remark: doses: 3.9 and 81 mg/rat/day (calculated by the author)
Result: There were no harmful effects on the fetuses with respect to external, skeletal and visceral anomalies; the stillborn litter number tended to increase; the postnatal development

of the off- spring was normal

25-APR-2001 (69)

Species: rat Sex:

Strain:

Route of admin.: oral feed

Exposure period: d 0 to 21 of gestation

Frequency of

treatment: daily

Duration of test:

Doses: 13 or 270 mg/kg/d

Control Group:

Method:

Year: GLP:

Test substance: other TS: N-oxydiethylene-2-benzothiazole sulfenamide (NOBS)

Result: no adverse effect on fetal or postnatal development

25-APR-2001 (70)

- 53/60 -

Species: Sex:

Strain:

Route of admin.:
Exposure period:
Frequency of
 treatment:
Duration of test:

Doses:

Control Group:

Method:

Year: GLP:

Test substance:

Remark: method: test compounds were tested for embryotoxicity and induction of malformations on three-day chicken embryos

Result: increased frequency of malformations

25-APR-2001 (71) (72)

5.10 Other Relevant Information

Type: Biochemical or cellular interactions

Remark: the conjugation between MBS and lysine, cysteine, and

glyzine was confirmed

Source: Bayer AG Leverkusen

20 - OCT - 1993 (73)

Type: other

Remark: Intratracheal injection of MBS powder caused pathological

alterations in the lungs (interstitial productive process,

emphysema, bronchitis)

Source: Bayer AG Leverkusen

07-APR-1992 (74)

Type:

Remark: Generation date of chap. 5: March 1992

Source: Bayer AG Leverkusen

20-OCT-1993

5.11 Experience with Human Exposure

26-MAY-1994

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Date: 15-OCT-2001
6. References ID: 102-77-2

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7.1 End Point Summary

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7.2 Hazard Summary

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7.3 Risk Assessment

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